# **PCT**

# WOORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



## INTERNATIONAL APPLICATIOON PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

		<del></del>
(51) International Patent Classification <sup>5</sup> : : C08F 220/28, 220/04 C09D 4/00, C08F 246/00	AI	(11) International Publication Number: WO 93/245 (43) International Publication Date: 9 December 1993 (09.12.
(21) International Application Number: PCT/US  (22) International Filing Date: 2.24 May 1993  (30) Priority data:  887,647 29 May 199292 (29.05.92)  887,673 29 May 199292 (29.05.92)  887,645 29 May 199292 (29.05.92)  887,644 29 May 199292 (29.05.92)  (71) Applicant: UNION CARBIDE CHHEMICALS  TICS TECHNOLOGY CORPORRATION [US Old Ridgebury Road, Danbury, CTT 06817-000)  (72) Inventors: JENKINS, Richard, Duantne; 309 Fletc  Hurrican, WV 25526 (US). BASSEET, David, Ro 2018 Huber Road, Charleston, WWV 25314 (US)  Gregory, Dean; 216 Kilbeck Drivive, Cary, N (US). SMITH, Danny, Elwood; 5324 Shad  Drive, Cross Lanes, WV 25313 (I(US). ARGY LOS, John, Nicholas; 35 Michaelel Street, Sco  WV 25560 (US). LOFFUS, Jamenes, Edward  Creek Crossing, Hurricane, WV 25:5526 (US).	(24.05.9)  & PLA //US]; I (US) her Lar obinsor ). SHA IC 275 dowbrod ROPO tt Depo	& Plastics Technology Corporation, 39 Old Ridgeb Road, Danbury, CT 06817-0001 (US).  (81) Designated States: AU, BB, BG, BR, CA, FI, HU, JP, I KR, LK, MG, MN, MW, NO, NZ, PL, RO, RU, S European patent (AT, BE, CH, DE, DK, ES, FR, C GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (I BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, T TG).  Published  With international search report.  Before the expiration of the time limit for amending claims and to be republished in the event of the receipt amendments.

(54) Title: COMPLEX HYDROPHOBE E COMPOUNDS, MACROMONOMERS AND MACROMONOMER-CONTAINING POLYMERS

#### (57) Abstract

This invention relates to polymers which comprise the reaction product of: (A) about 0-99.9 weight percent of one or more nonionic, cationic, anionic or amphoteric c monomers; (B) about 0-99.9 weight percent of one or more monoethylenically unsaturated monomers different from (A); (C) alabout 0.1-100 weight percent of one or more monoethylenically unsaturated monomers different from (A) and (B); and (L(D) about 0-20 weight percent or greater of one or more polyethylenically unsaturated monomers different from (A), (B) and (C).). When the polymers are added to latex paints and the pH adjusted, the viscosity of the paint is increased, brush drag is increased, and the paint rheology is otherwise improved. This invention also relates to macromonomer compounds useful as copolymeriziable monomers in various polymerization methods and as emulsifiers in the emulsion or suspension polymerization of various n monomers to produce an aqueous suspension of polymer particles. This invention further relates to complex hydrophobe compounded and to alkoxylation products, i.e., condensation reaction products of alkylene oxides and complex hydrophobe compounded having at least one active hydrogen and to alkoxylation products.

# FOOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify Statates party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	FR	France	MR	Mauritania
AU	Australia	GA	Gabon	MW	Malawi
88	Barbados	CB	United Kingdom	NL	Netherlands
BE	Belgium	GN	Guinca	NO	Norway
BF	Burkina Faso	GR	Greece	ΉZ	New Zealand
BG	Bulgaria	HU	Hungary	PŁ	Poland
BJ	Benin	IE	Ireland	PT	Portugal
BR	Brazil	IT	ltaly	RO	Romania
CA	Canada	JP	Japan	RU	Russian Federation
CF	Central African Republic	KP	Democratic People's Republic	SD	Sudan .
CG	Congo		of Korea	SE	Sweden
CH	Switzerland	KR	Republic of Korea	SK	Slovak Republic
Ċı	Côte d'Ivoire	KZ	Kazakhstan	SN	Senegal
СМ	Cameroon	1.1	Liechtenstein	รบ	Soviet Union
CS	Czechoslovakia ·	LK	Sri Lanka	TĐ	Chad
CZ	Czech Republic	1.U	Luxembourg	TG	Togo
DE	Germany	MC	Monaco-	UA	Ukraine
DK	Denmark	MG	Madagascar	us	United States of America
ES	Spain	MI.	Mali	VN	Viet Nam
Pi	Finland	MN	Mongolia		

- 1 -

# COMPLEX HYYDROPHOBE COMPOUNDS, MACROMONOMERS AND MAACROMONOMER-CONTAINING POLYMERS

### Brief Summary of the Invention

## Technical Field !

This invention relates to polymers which contain macromonomers and which can contain nonionic, cationic, anionic and/or amphoteeric monomers. The polymers are especially useful as thickeners for acqueous systems, e.g., certain of the polymers are soluble in, or swwelled by, an aqueous alkaline medium to provide thickeners for use in aqueous coating compositions, especially latex paints. This invention also relates to macromonomers containing complex hydrophobic groups and to complex hydrophobe compounds.

#### Background of tithe Invention

Thickeners for aqueous systems are needed for various purposes, such as for architectual coatings, industrial coatings, automotive coatitings and the like to improve rheology of the coatings. Hydroxyethyl ceellulose is a well known thickener for aqueous systems, but it has various deficiencies in that excessive amounts must be used annut the rheology of the thickened system is inadequate. Various ethoxyldated carboxyl-functional polymers which form alkali soluble thickeneers are also known, but these have various deficiencies, including inadequate hydrolytic stability.

It has long been desired to provide superior thickeners for aqueous systems which are highly efficient, which better resist hydrolysis, and \ which provide better rheology. This is achieved herein by providing new polymers which possess these desired characteristics.

#### Disclosure of thhe Invention

This invention relates in part to polymers (I) comprising the reaction prooduct of:

・メレーさい。 ニャリ・・・

- (A) ) about 0-99.9, preferably about 10-70, weight percent of one or more nonnionic, cationic, anionic and/or amphoteric monomers;
- (B) ) about 0-99.9, preferably about 30-85, weight percent of one or more monoethylenically unsaturated monomers, typically ethyl acrylate, ddifferent from component (A);
- (C)) about 0.1-100, preferably about 5-60, weight percent of one or more monoethylenically unsaturated macromonomers different from exomponents (A) and (B); and
- (D)) about 0-20, preferably about 0-10, weight percent or greater of one our more polyethylenically unsaturated monomers, typically trimethylol propane triacrylate, different from components (A), (B) and (C).).

Thiris invention also relates in part to an emulsion of the above-identifiedd polymer (I) in water, which emulsion is useful as a thickening agennt in aqueous compositions. In order to obtain the thickening effect, the polymer (I) is dissolved in the aqueous composition to bbe thickened.

Thins invention further relates in part to an aqueous composition, and more particularly an improved latex paint composition countaining the above-defined polymer (I).

Thins invention yet further relates in part to a process for thickening an aaqueous composition which comprises adding the above-defined poolymer (I) to an aqueous composition and dissolving the polymer (I) in the aqueous composition.

TThis invention relates in part to polymers (II) comprising the reaction prooduct of:

(AA) about 1-99.9, preferably about 10-70, weight percent of one or more alppha, beta-monoethylenically unsaturated carboxylic acids, typically methacrylic acid;

550 33: ----

(BB) about 0-98.9, preferably about 30-85, weight percent of one or more monnoethylenically unsaturated monomers, typically ethyl acrylate, different from component (A);

(CC) about 0.1-99, preferably about 5-60, weight percent of one or more monoethylenically unsaturated macromonomers different from components (A) 2 and (B); and

(DD) about 0-20, preferably about 0-10, weight percent or greater of one orr more polyethylenically unsaturated monomers, typically trimethylol proppane triacrylate, diffferent from components (A), (B) and (C).

This invention also relates in part to an emulsion of the above-identified ! polymer (II) in water, which emulsion is useful as a thickening agent in aqueous compositions. In order to obtain the thickening effect:t, the polymer (II) is dissolved in the aqueous composition to be thickened.

This invention further relates in part to an aqueous composition, andd more particularly an improved latex paint composition containing the abbove-defined polymer (II).

This invention yet further relates in part to a process for thickening an acqueous composition which comprises adding the above-defined poolymer (II) to an aqueous composition and dissolving the polymer (II) in the aqueous composition.

This invention relates in part to polymers (III) comprising the reaction product of:

(AA) from 1 to about 99.9 weight percent, preferably from about 60 to 3 95 weight percent and more preferably from about 70 to 95 weight percent, of one or more nonionic monoethylenically unsaturated mornomers, for example, acrylamide, N,N-dimethyl acrylamide, vinyyl pyrrolidone and hydroxyethyl acrylate; and

(BB) from about 0.1 weight percent to 99 weight percent, preferably from about 5 to 40 weight percent and more preferably from ε about 5 to 30 weight percent, of one or more

monoethylenicably unsaturated macromonomers different from component (A).

This invention also relates in part to an emulsion of the above-identitified polymer (III) in water, which emulsion is useful as a thickening agent in aqueous compositions. In order to obtain the thickening effects, the polymer (III) is dissolved in the aqueous composition to bbe thickened.

TThis invention further relates in part to an aqueous composition, and more particularly an improved latex paint composition conntaining the above-defined polymer (III).

Thins invention yet further relates in part to a process for thickening an acqueous composition which comprises adding the above-defined poolymer (III) to an aqueous composition and dissolving the polymer (IIIII) in the aqueous composition.

TThis invention relates to compounds represented by the formula:

$$R^4$$
 | R1-(OR<sup>2</sup>)<sub>z</sub>-R<sup>3</sup>-(-C=CR<sup>5</sup>R<sup>6</sup> (I)

wherein:

RR<sup>1</sup> is a monovalent residue of a substituted or unsubstituted ecomplex hydrophobe compound;

exach  $R^2$  is the same or different and is a substituted or unsubstituted ddivalent hydrocarbon residue;

RR<sup>3</sup> is a substituted or unsubstituted divalent hydrocarbon ressidue;

 ${\rm RR}^4, {\rm R}^5$  and  ${\rm R}^6$  are the same or different and are hydrogen or a substituted or unsubstituted monovalent hydrocarbon residue; and

zzis a value of 0 or greater.

- 5 -

Thhis invention relates to compounds represented by the formula:

$$R_{1}$$
-(O-CH<sub>2</sub>)<sub>a i</sub>

$$R_{3}$$
-(OR<sub>4</sub>)<sub>x</sub>-(OR<sub>5</sub>)<sub>y</sub>-OR<sub>6</sub>

$$R_{2}$$
-(O-CH<sub>2</sub>)<sub>b</sub>,

(i)

wherein  $R_1$  and  $l R_2$  are the same or different and are hydrogen or a substituted or unnsubstituted monovalent hydrocarbon residue, R3 is a substituted or unnsubstituted divalent or trivalent hydrocarbon residue, each R44 is the same or different and is a substituted or unsubstituted dilivalent hydrocarbon residue, each  $R_5$  is the same or different and is a a substituted or unsubstituted divalent hydrocarbon residue, R<sub>6</sub> is hyydrogen, a substituted or unsubstituted monovalent hydrocarbon resisidue or an ionic substituent, a and b are the same or different and aree a value of 0 or 1, and x and y are the same or different and are a value of 0 or greater; provided (i) at least two of  $R_1$ ,  $R_2$ ,  $R_3$ ,  $R_4$ ,  $R_5$  aand  $R_6$  are a hydrocarbon residue having greater than 2 carbon atoms iiin the case of  $R_1$ ,  $R_2$  and  $R_6$  or having greater than 2 pendant carbon  $\epsilon$  atoms in the case of  $R_3$ ,  $R_4$  and  $R_5$ , (ii) at least one of  $R_4$  and  $R_5$  is othher than -CH<sub>2</sub>CH<sub>2</sub>- when  $R_1$  and  $R_2$  are nonylphenyl, R<sub>3</sub> is CH-, R<sub>6</sub> is s hydrogen or SO<sub>3</sub>, a is a value of 1, b is a value of 1, and at least one  $\epsilon$  of x and y is a value of 1 or greater, and (iii)  $R_{\lambda}$  is -CH<sub>2</sub>CH<sub>2</sub>- and xx is a value of at least 2 when  $R_1$  is hexadecyl,  $R_2$  is tetradecyl, R<sub>3</sub> is CH-, at least one R<sub>5</sub> is -CH<sub>2</sub>CH(tetradecyl)-, R<sub>6</sub> is hydrogen, y is a 1 value of 1 or greater, a is a value of 1, and b is a value of 0.

440 Surmouth

TThis invention also relates to compounds represented by the formula::

$$\begin{array}{c} \text{R}_{7}\text{-}(\text{OCH}_{2})_{d}\text{-}\text{R}_{99}\text{-}(\text{OR}_{10})_{f}\text{-}\text{OR}_{11} \\ \\ \text{R}_{1:15} \\ \text{I} \\ \\ \text{R}_{8}\text{-}(\text{OCH}_{2})_{e}\text{-}\text{R}_{122}\text{-}(\text{OR}_{13})_{g}\text{-}\text{OR}_{14} \end{array} \tag{ii)}$$

wherein  $R_7$  and d $R_8$  are the same or different and are hydrogen or a substituted or vunsubstituted monovalent hydrocarbon residue,  $R_{11}$ and  $R_{14}$  are those same or different and are hydrogen, a substituted or unsubstituted rmonovalent hydrocarbon residue or an ionic substituent,  $R_{99}$  and  $R_{12}$  are the same or different and are a substituted or vunsubstituted divalent or trivalent hydrocarbon residue, each RR<sub>10</sub> is the same or different and is a substituted or unsubstituted ddivalent hydrocarbon residue, each  $R_{13}$  is the same or different and is a substituted or unsubstituted divalent hydrocarbon residue, R<sub>15</sub> is 3 a substituted or unsubstituted divalent hydrocarbon residue, d and e e are the same or different and are a value of 0 or 1. and f and g aree the same or different and are a value of 0 or greater; provided at leasist two of  $R_7$ ,  $R_8$ ,  $R_9$ ,  $R_{10}$ ,  $R_{11}$ ,  $R_{12}$ ,  $R_{13}$ ,  $R_{14}$  and  $R_{15}$ are a hydrocarbon residue having greater than 2 carbon atoms in the case of R<sub>7</sub>, R<sub>8</sub>, l R<sub>11</sub> and R<sub>14</sub> or having greater than 2 pendant carbon atoms in the cause of  $R_9$ ,  $R_{10}$ ,  $R_{12}$ ,  $R_{13}$  and  $R_{15}$ .

Thhis invention further relates to alkoxylation product mixtures whichh can have relatively narrow alkoxylation product distributions. These product mixtures are relatively free from large amounts of subbstantially higher alkoxylation moieties, i.e, those having at least t three more alkoxyl groups than the average peak

-7-

alkoxylate speciee. Advantageously, these narrow distributions can be obtained where t the most prevalent alkoxylation moiety has four or greater alkoxy uunits, that is, in the regions which generally have a relatively wide rrange of alkoxylation species.

#### **Detailed Description**

Illustrative nonionic, cationic, anionic and amphoteric monomers usefuld in this invention include those monomers which impart water solubility to the polymer. Preferably, a large proportion of component (A), is employed to impart water solubility to the polymers of this i invention. The key to water solubility lies in positioning sufficient numbers of hydrophilic functional groups along the backboone or side chains. Suitable functional groups which impart water solubility and suitable nonionic, cationic, anionic and amphoteric monomers useful in this invention are described in Water-Soluble Pcolymers, Synthesis, Solution Properties and Applications, ACCS Symposium Series 467, American Chemical Society (1991), wwhich is incorporated herein by reference. Mixtures of nonionic, cationic, anionic and amphoteric monomers may be employed in thiss invention, e.g., mixtures of nonionic monomers, mixtures of noninionic and cationic monomers, etc.

Illustrative nonionic monomers useful in this invention include, for example, acrylamide, N, N-dimethyl acrylamide, vinyl pyrrolidone, ethyylene oxide, vinyl alcohol, vinyl acetate, N-vinylpyrrolidinonne, hydroxyethyl acrylate and the like including mixtures thereoff. Illustrative cationic monomers useful in this invention includde, for example, ammonium, sulfonium and phosphonium salts, preferably quarternary ammonium salts such as diallyldimethylammonium chloride, diallyldiethylammonium chloride, diethylaminoethyl methacrylate, dimethylaminoethyl methacrylate, mnethacryloyloxyethyltrimethylammonium sulfate, methacryloyloxyethyltrimethylammonium chloride, 3-

(methacrylamiddo)propyltrimethylammonium chloride and the like including mixtuures thereof.

Illustrative anionic monomers useful in this invention include, for exaample, acrylic acid, methacrylic acid, maleic anhydride, p-stytyrene carboxylic acids, p-styrene sulfonic acids, vinyl sulfonic acid, 2-2-sulfoethyl methacrylate, 3-sulfopropyl methacrylate, 3-sulfopropyl accrylate, 2-acrylamido-2-methylpropane sulfonic acid. 3-acrylamido-3-1-methylbutanoic acid and the like including mixtures thereof and saltits thereof. Illustrative amphoteric monomers useful in this invention contain zwitterions on the same monomers, i.e., betaines, or along the same polymer backbone, i.e., ampholytes, and include, for exaample, N,N-dimethyl-N-methacryloxyethyl-N-(3sulfopropyl)amnmonium-betain, N,N-dimethyl-Nmethacrylamiddopropyl-N-(3-sulfopropyl)ammonium-betain, 1-(3sulfopropyl)-2-vvinylpyridinium-betain, 3-(2-acrylamido-2-methylpropyldimethyl:lammonio)-1-propanesulfonate, N-vinylpyrrolidone-co-N,N-dimethyl-NN-methacroyloxyethylammoniopropanesulfonate, Nvinylpyrrolidonae- co-N,N-dimethyl-Nmethacroylamiddopropylammonio- propanesulfonate, Nvinylpyrrolidonae-co-2-vinylpyridinio-propanesulfonate and the like including mixtures thereof.

Preeferably, a large proportion of one or more alpha, betamonoethylenicaally unsaturated carboxylic acid monomers can be present in the ppolymers of this invention. Various carboxylic acid monomers can I be used, such as acrylic acid, methacrylic acid, ethacrylic acid,I, alpha-chloroacrylic acid, crotonic acid, fumaric acid, citraconic acid,, mesaconic acid, itaconic acid, maleic acid and the like including nmixtures thereof. Methacrylic acid is preferred. A large proportion of carboxylic acid monomer is essential to provide a polymeric struccture which will solubilize and provide a thickener when reacted wwith an alkali like sodium hydroxide. Thee polymers of this invention can also contain a significant propoortion of one or more monoethylenically unsaturated monomers. Thee preferred monomers provide water insoluble polymers when I homopolymerized and are illustrated by acrylate and methacrylate eststers, such as ethyl acrylate, butyl acrylate or the corresponding mmethacrylate. Other monomers which can be used are styrene, alkyl styrenes, vinyl toluene, vinyl acetate, vinyl alcohol, acrylonitrile, vinnylidene chloride, vinyl ketones and the like.

Nonreactive modnomers are preferred, those being monomers in which the single ethylenic group is the only group reactive under the conditions of poblymerization. However, monomers which include groups reactive 1 under baking conditions or with divalent metal ions such as zinc oxide may be used in some situations, like hydroxyethyl acrylate.

Othher illustrative monoethylenically unsaturated monomers usefuul in this invention include, for example, propyl methacrylate, is sopropyl methacrylate, butyl methacrylate, n-amyl methacrylate, seec-amyl methacrylate, hexyl methacrylate, lauryl methacrylate, statearyl methacrylate, ethyl hexyl methacrylate, crotyl methacrylate, cirinnamyl methacrylate, oleyl methacrylate, ricinoleyl methacrylate, hyydroxy ethyl methacrylate, hydroxy propyl methacrylate, vinnyl propionate, vinyl butyrate, vinyl tert-butyrate, vinyl caprate, vininyl stearate, vinyl laurate, vinyl oleate, vinyl methyl ether, vinyl ethyvl ether, vinyl n-propyl ether, vinyl iso-propyl ether, vinyl n-butyl ethher, vinyl iso-butyl ether, vinyl iso-octyl ether, vinyl phenyl ether, a-c-chlorovinyl phenyl ether, vinyl /-naphthyl ether, methacryonitrilde, acrylamide, methacrylamide, N-alkyl acrylamides, N-2-aryl acrylamides, N-vinyl pyrrolidone, N-vinyl-3morpholinones, I N-vinyl-oxazolidone, N-vinyl-imidazole and the like including mixtuures thererof.

Thee macromonomers useful in this invention can be represented by the formula (I) above. The macromonomer

compounds useful in this invention can be prepared by a number of conventional processes, except for inclusion of the complex hydrophobe compounds described herein. Illustrative processes are described, for example, in U.S. Patent Nos. 4,514,552, 4,600,761, 4,569,965, 4,384,4,096, 4,268,641, 4,138,381, 3,894,980, 3,896,161, 3,652,497, 4,509,949, 4,2265,754, 3,915,921, 3,940,351, 3,035,004, 4,429,097, 4,421,902, 4,167,502, 4,764,4,554, 4,616,074, 4,464,524, 3,657,175, 4,008,202, 3,190,925, 3,794,608, 4,3388,239, 4,939,283 and 3,499,876. Other macromonomer compounds whitch may be useful in this invention include complex hydrophobe-conntaining oligomers dislosed in copending U.S. Patent Application Seririal No. 07/887,646, filed May 29, 1992, which is incorporated heerein by reference.

I In a preferred embodiment, the macromonomer compounds of this invention can be prepared by coupling a compound represented by y the formula:

$$R^{1}$$
- $(OR^{2})_{z}$ -OH (II)

with an ethyleenically unsaturated compound such as alpha, alphadimethyl meta i isopropenyl benzyl isocyanate having the formula:

$$H_{2} \circ C = C \qquad CH_{3} \qquad CH_{3} \qquad CH_{3} \qquad CH_{3} \qquad CH_{3} \qquad (III)$$

wherein  $R^1$ ,  $R^2$  and z are as defined above. This coupling reaction can be carried l out in the presence of a catalyst such as bismuth octoate, dibutyly tin dilaurate and the like. The coupling reaction can

be conducted at a a temperature of about 50°C to about 150°C and a pressure of about 1 atmosphere or so. Stoichiometric amounts of starting materiaals can generally be employed.

It is understood that surfactants or alcohols with more than one hydroxxyl, or compounds that have amine functionality, and the like can be uused. Other unsaturated mono-, di-, and triisocyanates, s such as isocyanato ethyl methacrylate, allyl isocyanate, and i the like can also be used. Other monoisocyanates useful in preparring macromonomer compounds of this invention have the formulula:

$$Z - C - O - Y - N = C = O$$
(IV)

where Y is a substituted or unsubstituted divalent hydrocarbon residue such as  $\epsilon$  defined for  $R^3$  herein, and Z is the residue of an ethylenically unnsaturated compound such as  $CH_2=CH_2$ ,  $CH_2=C(CH_3)$ - onr  $CH_2=CH-CH_2$ -.

In another embodiment, the compound represented by formula (II) cann react with an alpha, beta-unsaturated dicarboxylic acid or acid anhhydride having the formula:

(V)

to produce an easter, wherein  $R^7$ ,  $R^8$ ,  $R^9$  and  $R^{10}$  are the same or different and arre hydrogen or a substituted or unsubstituted hydrocarbon ressidue, preferably an alkyl residue. If  $R^7$ ,  $R^8$ ,  $R^9$  and  $R^{10}$  are hydrogen, then the acrylic ester is formed; if  $R^7$  and  $R^{10}$  are hydrogen and  $R^8$  and  $R^9$  are methyl, then the methacrylic ester is formed; if  $R^7$  annd  $R^{10}$  are methyl and  $R^8$  and  $R^9$  are hydrogen, then the crotonic ester is formed. Other anhydrides, such as itaconic anhydride, crottonic anhydride, allyl succinic, methylene malonic, and the like cann also be used.

TThis reaction can be carried out in the presence of a catalyst such ass pyridine, triethylamine and the like. The reaction can be conducteed at a temperature of about ambient to about 120°C and a pressure  $\epsilon$  of about 1 atmosphere or so. Stoichiometric amounts of starting matterials can generally be employed.

In addition, cyclic alpha, beta-unsaturated dicarboxylic acidid anhydrides, such as maleic anhydride can also be used:

(VI)

wherein R<sup>11</sup> annd R<sup>12</sup> are the same or different and are hydrogen or a substituted or uunsubstituted hydrocarbon residue, preferably halogen, cyanoggen, hydroxyl, lactam and lactone groups, aryl,

aralkyl, substituuted alkyl, aryl or aralkyl groups, such as methyl, ethyl, propyl, oct.tyl, decyl, phenyl, tolyl, xylyl, benzyl, cyclohexyl, and the like, or -SO<sub>33</sub>H. Examples of such compounds are chloromaleic anhydride, citraaconic anhydride, fumaric anhydride, mescaconic anhydride, phennyl maleic anhydride, benzyl maleic anhydride, sulfomaleic anhydride, aconitic anhydride, and the like.

Other synthetic routes such as transesterification or nucleophilic substitution reactions can be used. For example, the compound represented by formula (II) can react with an acid halide having the formula:

wherein R<sup>13</sup> is t the residue of an ethylenically unsaturated compound such as CH<sub>2</sub>=CHH-, CH<sub>2</sub>=C(CH<sub>3</sub>)- or CH<sub>2</sub>=CH-CH<sub>2</sub>-, and X is NH<sub>2</sub> or halogen such as ; chloro, bromo, fluoro or iodo. This reaction can be carried out in thoe presence of a catalyst such as pyridine, triethylamine annd the like. The reaction can be conducted at a temperature of about ambient to about 120°C and a pressure of about 1 atmosphere or sco. Stoichiometric amounts of starting materials can generally be empployed.

Inn yet another embodiment, the compound represented by formula (II) can react with an epoxide having the formula:

$$R^{14}$$
  $O$   $CH_2$   $CH_2$  (VIII)

wherein  $R^{14}$  is a substituted or unsubstituted hydrocarbon residue, preferably the residue of an ethylenically unsaturated compound such as  $CH_2$ =CFH-,  $CH_2$ = $C(CH_3)$ - or  $CH_2$ =CH- $CH_2$ -. If  $R^{14}$  is not a reactive substitutent, then the resulting product having the formula:

$$R^{1}$$
- $(OR^{2})_{z}$ -O-4-CH-CH $_{2}$ -O- $R^{14}$  (IX)

can be reacted with an unsaturated hydrocarbon compound such as an isocyanate hhaving the formula:

$$R^{15}-N=C=O (X)$$

to provide a maacromonomer compound having the formula:

$$RR^{1}$$
— $(OR^{2})_{z}$ — $O$ — $CH$ — $CH_{2}$ — $O$ — $R^{14}$ 
 $O$ 
 $O$ = $C$ 
 $NH$ 
 $15$ 

(XI)

wherein  $R^1$ ,  $R^2$ ,  $R^{14}$  and z are as defined above and  $R^{15}$  is the residue of an ethylenicaally unsaturated compound such as  $CH_2$ =CH-,

CH<sub>2</sub>=C(CH<sub>3</sub>)- onr CH<sub>2</sub>=CH-CH<sub>2</sub>-. This reaction can be carried out in the presence of an catalyst such as bismuth octoate, dibutyl tin dilaurate and the like. The reaction can conducted at a temperature of about ambient to about 120°C and a pressure of about 1 atmosphere or so. Stoichiomnetric amounts of starting materials can generally be employed.

The macromonomer can be prepared by a neat process (without solvent), or in solution. When neat, the complex hydrophobe compound having at least one active hydrogen such as a surfactant or poly(oxyethyldene) derivative can be melted and sparged with nitrogen before c charging the reactants to remove moisture; when in solution, water isis removed by way of an azeotrope. Then the catalyst and capping matterials are charged, and the reaction mixture heated until the reaction goes to completion. The reaction stoichiometry of the surfactant annd the capping material may be optimized to promote thickener efficiency and minimize coagulum formation during the subsequent polyumerization. Afterwards, water can be added to the neat macromonomer to lower its viscosity. The material made in solution is recovered as a solid after solvent removal, or can be put into aqueous solution by the addition of water during azeotropic distillation.

Th'he unsaturation portion of the macromonomer compounds of thhis invention can be derived from olefinically unsaturated compounds containing at least one activated carbon-to-carbon olefinic ddouble bond which readily functions in polymerization oof the macromonomer compound. Suitable olefinically unsaaturated compounds include, for example, acrylic acids typified by a acrylic acid itself, methacrylic acid, ethacrylic acid, alpha-chloro-acryylic acid, alpha-cyano acrylic acid, beta-methyl acrylic acid (crotstonic acid), alpha-phenyl acrylic acid, beta-acryloxy propionic acid, ssorbic acid, alpha-chloro sorbic acid, angelic acid, cinnamic acid, pp-chloro cinnamic acid, beta-styryl acrylic acid (1-

WO 93/24544 PCT/US93/04872

- 16 -

carboxy-4-phenyyl butadiene-1,3), itaconic acid, citraconic acid, mesaconic acid,!, glutaconic acid, aconitic acid, maleic acid, fumaric acid, and tri-canrboxy ethylene. Also included are polycarboxylic acids and those e acid anhydrides, such as maleic anhydride, wherein the anhydride ggroup is formed by the elimination of one molecule of water from two o carboxyl groups located on the same polycarboxylic acid molecule.

Suitable acid anhydrides include alpha, betaunsaturated diccarboxylic acid anhydrides such as maleic anhydride, chloromaleic annhydride, citraconic anhydride (methyl maleic), fumaric anhydride, phenyl maleic anhydride, benzyl maleic anhydride, sulfdomaleic anhydride, aconitic anhydride and the like. Other unsaturated anhydrides useful in this invention include itaconic anhydride, methylene malonic anhydride, allyl succinic anhydride and t the like.

Illustrative substituted and unsubstituted divalent hydrocarbon ressidues represented by R<sup>2</sup> in formula (I) above include those described I for the same type of substituents in formulae (i) and (ii) below. Illustrative substituted and unsubstituted monovalent hydrocarbon ressidues represented by R<sup>4</sup>, R<sup>5</sup> and R<sup>6</sup> in formula (I) above include thhose described for the same type of substituents in formulae (i) andd (ii) below.

Illustrative R<sup>3</sup> substituents include, for example, the organic residuee of ethers, esters, urethanes, amides, ureas, urethanes, anhydrides and the like including mixtures thereof. The R<sup>3</sup> substituent can be generally described as a "linkage" between the complex hydrophhobe bearing surfactant or alcohol, and the unsaturation poortion of the macromonomer compound. Preferred linkages include the following: urethane linkages from the reaction of an isocyanate with a nonionic surfactant; urea linkages from the reaction of an isisocyanate with an amine bearing surfactant; unsaturated estiters of surfactants such as the esterification product of

- 17 -

a surfactant with of an unsaturated carboxylic acid or an unsaturated anhhydride; unsaturated esters of alcohols; esters of ethyl acrylate oligometers, acrylic acid oligometers, and allyl containing oligometers; half elesters of surfactants such as those made by the reaction of a surrfactant with maleic anhydride; unsaturated ethers prepared by reaccting vinyl benzyl chloride and a surfactant or by reacting an allylil glycidyl ether with a surfactant, alcohol, or carboxylic acid.

Thee oxyalkylene moieties included in the macromonomer  $\epsilon$  compounds of formula (I) may be homopolymers or block or random a copolymers of straight or branched alkylene oxides. Mixtures of alkylene oxides such as ethylene oxide and propylene oxide may be employed. It is understood that each  $R^2$  group in a particular substitutent for all positive values of z can be the same or different.

Thee complex hydrophobe compounds having at least one active hydrogen i useful in preparing the macromonomer compounds useful in this invvention can be represented by the formula:

$$R_{1}$$
-(O-CH<sub>2</sub>)<sub>a 1</sub>

$$R_{3}$$
-(OR<sub>4</sub>)<sub>x</sub>-(OR<sub>5</sub>)<sub>y</sub>-OR<sub>6</sub>

$$R_{2}$$
-(O-CH<sub>2</sub>)<sub>b</sub>,

(i)

wherein  $R_1$  and l  $R_2$  are the same or different and are hydrogen or a substituted or unnsubstituted monovalent hydrocarbon residue,  $R_3$  is a substituted or unnsubstituted divalent or trivalent hydrocarbon residue, each  $R_4$  is the same or different and is a substituted or unsubstituted divalent hydrocaarbon residue, each  $R_5$  is the same or different and is a

substituted or unsubstituted divalent hydrocarbon residue, R6 is hydrogen, a substituted or unsubstituted monovalent hydrocarbon residue or an ionic subsstituent, a and b are the same or different and are a value of 0 or 1, and x  $\epsilon$  and y are the same or different and are a value of 0 or greater; provided at least two of R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub>, R<sub>5</sub> and R<sub>6</sub> are a hydrocarbon ressidue having greater than 2 carbon atoms in the case of  $R_1$ ,  $R_2$  and  $R_6$ ; or having greater than 2 pendant carbon atoms in the case of R<sub>3</sub>, R<sub>4</sub> and RR<sub>5</sub>. For purposes of the complex hydrophobes of formula (i) above, at least cone of R<sub>4</sub> and R<sub>5</sub> is other than -CH<sub>2</sub>CH<sub>2</sub>- when R<sub>1</sub> and R<sub>2</sub> are nonylphenyyl, R<sub>3</sub> is CH-, R<sub>6</sub> is hydrogen or SO<sub>3</sub>, a is a value of 1, b is a value of 1, and I at least one of x and y is a value of 1 or greater. Also for purposes of thee complex hydrophobes of formula (i) above, R<sub>4</sub> is -CH<sub>2</sub>CH<sub>2</sub>and x is a value of at least 2 when  $R_1$  is hexadecyl,  $R_2$  is tetradecyl,  $R_3$  is CH-, at least opne  $R_5$  is -CH<sub>2</sub>CH(tetradecyl)-,  $R_6$  is hydrogen, y is a value of 1 or greater, a i is a value of 1, and b is a value of 0. For purposes of the homopolymers 3 and macromonomers of formula (I) above, when z is a value of 0 and  $1 R^1$  is the residue of a complex hydrophobe of formula (i) in which  $R_1$  is hexadecyl, a is a value of 1,  $R_2$  is tetradecyl, b is a value of 0, R<sub>3</sub> is CH-, R<sub>4</sub> i is -CH<sub>2</sub>CH(tetradecyl)-, x is a value of 1, R<sub>5</sub> is -CH<sub>2</sub>CH<sub>2</sub>-, y is a value of 344 and  $R_6$  is hydrogen, then the  $-R^3(R^4)C = CR^5R^6$  portion of the macromonomer is other than the residue of maleic anhydride. Also for purposes off the homopolymers and macromonomers of formula (I) above, when  $R^{12}$  and  $R^3$  are hydrogen, z is a value of 34 and  $R^1$  is the residue of a complex hydrophobe of formula (i) in which R<sub>1</sub> is hexadecyl, a is a value of 1, 1 R2 is tetradecyl, b is a value of 0, R3 is CH-, R4 is -CH<sub>2</sub>CH(tetraddecyl)-, x is a value of 1, y is a value of 0 and R<sub>6</sub> is hydrogen, then the  $-R^3$ - $(RR^4)$ C= $CR^5R^6$  portion of the macromonomer is other than the residue of maldeic anhydride. For purposes of polymers (I) and (II) above, when z is a value of 0 and  $R^1$  is the residue of a complex hydrophobe of formula (i) in wwhich R<sub>1</sub> is hexadecyl, a is a value of 1, R<sub>2</sub> is tetradecyl, b is a value of 0, R33 is CH-, R4 is -CH2CH(tetradecyl)-, x is a value of 1, R5 is -CH<sub>2</sub>CH<sub>2</sub>-, y isis a value of 34, R<sub>6</sub> is hydrogen, and the -R<sup>3</sup>-(R<sup>4</sup>)C=CR<sup>5</sup>R<sup>6</sup>

WO 93/24544

PCT/US93/04872

- 19 -

portion of the maacromonomer is the residue of maleic anhydride, then the polymers of this innvention are other than a terpolymer of said macromonomer, s styrene and maleic anhydride. Also for purposes of polymers (I) and ((II) above, when  $R^2$  is -CH<sub>2</sub>CH<sub>2</sub>-, z is a value of 34 and  $R^1$  is the residue  $\epsilon$  of a complex hydrophobe of formula (i) in which  $R_1$  is hexadecyl, a is a vvalue of 1,  $R_2$  is tetradecyl, b is a value of 0,  $R_3$  is CH-,  $R_4$  is -CH<sub>2</sub>CH(tetraddecyl)-, x is a value of 1, y is a value of 0,  $R_6$  is hydrogen and the - $R^3$ -( $R^4$ )OC=CR $^5$ R $^6$  portion of the macromonomer is the residue of maleic anhydride, then the polymers of this invention are other than a terpolymer of said macromonomer, styrene and maleic anhydride.

Other complex hydrophobe compounds having at least one active hydropgen useful in preparing the macromonomer compounds usefull in this invention can be represented by the formula:

wherein  $R_7$  and l  $R_8$  are the same or different and are hydrogen or a substituted or unnsubstituted monovalent hydrocarbon residue,  $R_{11}$  and  $R_{14}$  are the  $\iota$  same or different and are hydrogen, a substituted or unsubstituted monovalent hydrocarbon residue or an ionic substituted,  $R_9$  and  $R_{12}$  are the same or different and are a substituted or unnsubstituted divalent or trivalent hydrocarbon residue, each  $R_{10}$  is the same or different and is a substituted or unsubstituted dilivalent hydrocarbon residue, each  $R_{13}$  is the same or different and is  $\epsilon$  a substituted or unsubstituted divalent hydrocarbon residue,  $R_{15}$  is an aubstituted or unsubstituted divalent hydrocarbon residue, d and  $\epsilon$  are the same or different and are a value of 0 or 1,

and f and g aree the same or different and are a value of 0 or greater; provided at leasest two of  $R_7$ ,  $R_8$ ,  $R_9$ ,  $R_{10}$ ,  $R_{11}$ ,  $R_{12}$ ,  $R_{13}$ ,  $R_{14}$  and  $R_{15}$  are a hydrocarbbon residue having greater than 2 carbon atoms in the case of  $R_7$ ,  $R_8$ ,  $R_{11}$  and  $R_{14}$  or having greater than 2 pendant carbon atoms in the case of  $R_9$ ,  $R_{10}$ ,  $R_{12}$ ,  $R_{13}$  and  $R_{15}$ .

Illustrative substituted and unsubstituted monovalent hydrocarbon ressidues contain from 1 to about 50 carbon atoms or greater and aree selected from alkyl radicals including linear or branched primaary, secondary or tertiary alkyl radicals, such as methyl, ethyl, nn-propyl, isopropyl, amyl, sec-amyl, t-amyl, 2-ethylhexyl and I the like; aryl radicals such as phenyl, naphthyl and the like; arylalkkyl radicals such as benzyl, phenylethyl, triphenylmethyletthane and the like; alkylaryl radicals such as octylphenyl, nopnylphenyl, dodecylphenyl, tolyl, xylyl and the like; and cycloalkyl radicals such as cyclopentyl, cyclohexyl, cyclohexylethyl and the like. The permissible hydrocarbon residues may contain fluorine, siliconn, or other non-carbon atoms.

Preferably, the substituted and unsubstituted hydrocarbon residues are selected from alkyl and aryl radicals which contain from abbout 1 to 30 carbon atoms or greater. More preferably, the alkyl radicals contain from 1 to 18 carbon atoms, while the aryl, arylalkyl, alkyldaryl and cycloalkyl radicals preferably contain from 6 to 18 carbon atoms or greater.

In a preferred embodiment of this invention,  $R_1$ ,  $R_2$ ,  $R_7$  and  $R_8$  can individually be a hydrocarbon radical represented by the formula:

$$R_{16}$$
-(OCH<sub>2</sub>)<sub>h</sub> \ R<sub>18</sub>- \ (iii) \ R<sub>17</sub>-(OCH<sub>2</sub>)<sub>i</sub>

- 21 -

wherein R<sub>16</sub> andd R<sub>17</sub> are as defined for R<sub>1</sub>, R<sub>2</sub>, R<sub>7</sub> and R<sub>8</sub> above, h and i are the same or different and are a value of 0 or 1, and R<sub>18</sub> is as defined for R<sub>3</sub> abbove. For compounds represented by formulae (i) and (ii), it is understated that each formula (iii) radical in a given compound may bbe the same or different and the R<sub>16</sub> and/or R<sub>17</sub> groups may then selves be a formula (iii) radical to provide complex hydrophobes of an dendritic or of a cascading nature as described below. Further, R<sub>4</sub>, R<sub>5</sub>, R<sub>10</sub> and R<sub>13</sub> can individually be a hydrocarbon raddical represented by the formula:

 $-CH[(OR_{19})_j)_jOR_{20}]$ 

(iv)

wherein  $R_{19}$  is any defined for  $R_4$ ,  $R_5$ ,  $R_{10}$  and  $R_{13}$  above,  $R_{20}$  is as defined for  $R_6$ ,  $RR_{11}$  and  $R_{14}$  above, and j is a value of 0 or greater.

Illuustrative ionic substituents for  $R_6$ ,  $R_{11}$ ,  $R_{14}$  and  $R_{20}$  include cationic: and anionic substituents such as sulfates, sulfonates, phospphates, carboxyls and the like.  $R_6$ ,  $R_{11}$ ,  $R_{14}$  and  $R_{20}$  may preferably t be an organic residue containing 1 or more hydroxyls or nitrogen derivivatives or epoxides or other reactive groups which may or may not t contain unsaturation.

Othher illustrative terminal groups which are described by  $R_6$ ,  $R_{11}$ ,  $R_{14}$  and and  $R_{20}$  include, for example, hydrocarbon residues which I may contain allylic or vinylic unsaturation, acrylic or methacrylic functionality, styryl or alpha-methylstyryl functionality, and the like, such as the reaction product between the terminal alcohol ( $R_6$ ,  $R_{11}$ ,  $R_{14}$  and  $R_{20} = H$ ) and glycidyl methacrylate, isocyanatoethyl | methacrylate, alpha, alpha-dimethyl-m-isopropenyl benzyl isocyanatte (m-TMI), and the like. Other examples of terminal groups may include hydrocarbon residues of alkyl, aryl, aralkyl, alkaryl, and cyclcloalkyl radicals which may or may not be substituted with one or morre of the following: hydroxyl, carboxyl, isocyanato,

amino, mono- o or disubstituted amino, quaternary ammonium, sulfate, sulfonaate, phosphate, epoxy, and the like and may or may not contain other nnon-carbon atoms including silicon or fluorine. Also included can bee divalent siloxy radicals. Other nonhydrocarbon terminal groupps may include sulfates, phosphates, and the like.

Illustrative divalent hydrocarbon residues represented by  $R_3$ ,  $R_4$ ,  $R_5$ ,  $RR_9$ ,  $R_{10}$ ,  $R_{12}$ ,  $R_{13}$ ,  $R_{15}$ ,  $R_{18}$  and  $R_{19}$  in the above formulae includde substituted and unsubstituted radicals selected from alkylene, -alkylene-oxy-alkylene-, -arylene-oxy-arylene-, arylene, alicyclilic radicals, phenylene, naphthylene, -phenylene- $(CH_2)_m(Q)_n(CEH_2)_m$ -phenylene- and -naphthylene-(CH<sub>2</sub>)<sub>m</sub>(Q)<sub>n</sub>(CEH<sub>2</sub>)<sub>m</sub>-naphthylene- radicals, wherein Q individually represents a substituted or unsubstituted divalent bridging group selected from -CCR<sub>21</sub>R<sub>22</sub>-, -O-, -S-, -NR<sub>23</sub>-, -SiR<sub>24</sub>R<sub>25</sub>- and -CO-, wherein  $R_{21}$  annd  $R_{22}$  individually represent a radical selected from hydrogen, alkyld of 1 to 12 carbon atoms, phenyl, tolyl and anisyl; R23, R<sub>24</sub> and R<sub>25</sub> inndividually represent a radical selected from hydrogen and methyl, and each m and n individually have a value of 0 or 1. More specific illlustrative divalent radicals represented by R<sub>3</sub>, R<sub>4</sub>, R<sub>5</sub>, R<sub>9</sub>, R<sub>10</sub>, R<sub>12</sub>, RR<sub>13</sub>, R<sub>15</sub>, R<sub>18</sub> and R<sub>19</sub> include, e.g., 1,1-methylene, 1,2ethylene, 1,3-ppropylene, 1,6-hexylene, 1,8-octylene, 1,12-dodecylene, 1,4-phenylene, 1,8-napthylene, 1,1'-biphenyl-2,2'-diyl, 1,1'-binaphthyl-2,2'-diyl, 2,2'-bisinaphthyl-1,1'-diyl and the like. The alkylene radicals may contain froom 2 to 12 carbon atoms or greater, while the arylene radicals may countain from 6 to 18 carbon atoms or greater. Preferably,  $R_3$ ,  $R_4$ ,  $R_5$ ,  $R_9$ ,  $R_{10}$ ,  $R_{12}$ ,  $R_{13}$ ,  $R_{15}$ ,  $R_{18}$  and  $R_{19}$  are an alkylene or aryylene radical. The permissible divalent hydrocarbon residues may c contain fluorine, silicon, or other non-carbon atoms.

Illilustrative trivalent hydrocarbon residues represented by  $R_3$ ,  $R_9$ ,  $R_{12}$ , and  $R_{18}$  in the above formulae includee substituted and unsubstituted radicals selected from CH-,  $C(R_{26})$ -,  $CR_{27}$ - and the

like, wherein R<sub>266</sub> is a substituted or unsubstituted monovalent hydrocarbon residue as described herein and R<sub>27</sub> is a substituted or unsubstituted divialent hydrocarbon residue as described herein.

Of course, it is to be further understood that the hydrocarbon residdues in the above formulae may also be substituted with any permissible substituent. Illustrative substituents include radicals containing from 1 to 18 carbon atoms such as alkyl, aryl, aralkyl, alkaryl aand cycloalkyl radicals; alkoxy radicals; silyl radicals such as --Si(R28)3 and -Si(OR28)3, amino radicals such as  $-N(R_{28})_2$ ; acyl raadicals such as  $-C(O)R_{28}$ ; acyloxy radicals such as -OC(O)R28; carboonyloxy radicals such as -COOR28; amido radicals such as -C(O)N(RR28)2 and -N(R28)COR28; sulfonyl radicals such as -SO<sub>2</sub>R<sub>28</sub>; sulfinylyl radicals such as -SO(R<sub>28</sub>)<sub>2</sub>; thionyl radicals such as  $-SR_{28}$ ; phosphhonyl radicals such as  $-P(O)(R_{28})_2$ ; as well as halogen, nitro, cyyano, trifluoromethyl and hydroxy radicals and the like, wherein eackh R28 can be a monovalent hydrocarbon radical such as alkyl, arryl, alkaryl, aralkyl and cycloalkyl radicals, with the provisos that in aamino substituents such as  $-N(R_{28})_2$ , each  $R_{28}$  taken together can also ocompromise a divalent bridging group that forms a heterocyclic radicial with the nitrogen atom, in amido substituents such as  $-C(O)N(RR_{28})_2$  and  $-N(R_{28})COR_{28}$ , each  $R_{28}$  bonded to N can also be hydrogena, and in phosphonyl substituents such as  $-P(O)(R_{28})_2$ , one  $R_{28}$  can by hydrogen. It is to be understood that each R<sub>28</sub> group in a poarticular substituent may be the same or different. Such hydrocarbonn substituent radicals could possibly in turn be substituted with a permissible substituent such as already herein outlined above.

Presserred alkylene oxides which can provide random or block oxyalkylenee units in the complex hydrophobe compounds represented by foformulae (i) and (ii) include alkylene oxides such as ethylene oxide, ppropylene oxide, 1,2-butylene oxide, 2,3-butylene oxide, 1,2- and 2,2,3-pentylene oxide, cyclohexylene oxide, 1,2-hexylene

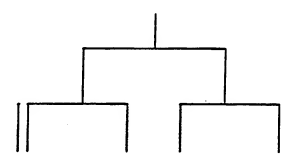
oxide, 1,2-octyleene oxide, 1,2-decylene oxide, and higher alpha-olefin epoxides; epoxiddized fatty alcohols such as epoxidized soybean fatty alcohols and epooxidized linseed fatty alcohols; aromatic epoxides such as styrene; oxide and 2-methylstyrene oxide; and hydroxy- and halogen-substituted alkylene oxides such as glycidol, epichlorohydrin and epibromohyydrin. The preferred alkylene oxides are ethylene oxide and propylylene oxide. Also included can be hydrocarbon residues from suubstituted and unsubstituted cyclic esters or ethers such as oxetane e and tetrahydrofuran. It is understood that the compounds represented by formulae (i) and (ii) herein can contain random and/or l block oxyalkylene units as well as mixtures of oxyalkylene unitits. It is further understood that each R<sub>4</sub>, R<sub>5</sub>, R<sub>10</sub>, R<sub>13</sub> and R<sub>19</sub> group in a particular substituent for all positive values of x, y, f, g and j j respectively can be the same or different.

Thee values of x, y, z, f, g and j are not narrowly critical and can vary over a wide range. For example, the values of x, y, z, f, g and j can rangge from 0 to about 200 or greater, preferably from about 0 to about 100 or greater, and more preferably from about 0 to about 50 or greater. Any desired amount of alkylene oxide can be employed, for exxample, from 0 to about 90 weight percent or greater based on the weeight of the complex hydrophobe compound.

Refrering to the general formulae (i) and (ii) above, it is appreciated that when R<sub>1</sub>, R<sub>2</sub>, R<sub>7</sub> and/or R<sub>8</sub> are a hydrocarbon residue of formulae (iii) above, the resulting compound may include any permissible e number and combination of hydrophobic groups of the dendritic or r cascading type. Such compounds included in the above general foormulae should be easily ascertainable by one skilled in the art.

In a preferred embodiment of this invention, the structure shown in formula (iii) can be a residue of the reaction product between epichlorohydrin and an alcohol, including those alcohols whose residues can be described by formula (iii), or a

phenolic, or a mnixture thereof. The structures which result can be described as complex hydrophobes of a dendritic or of a cascading nature. Pictoriaally, they can be described as shown below:



Illlustrative starter components useful for making the complex hydrophhobe compounds of this invention include those compounds haviring active hydrogens such as alcohols (mono-, di- and polyhydric alcohhols), phenols, carboxylic acids (mono-, di- and polyacids), and a amines (primary and secondary). Frequently, the organic compounnds contain up to about 100 or 150 carbons (in the case of polyol poolymers) and can contain aliphatic and/or aromatic structures. Mostst often, the starter compounds are selected from the group of mono-, c di- and trihydric alcohols having up to about 30 carbon atoms or r greater. The organic compounds having active hydrogens can boe the product of hydroformylation/hydrogenation reactions. Other useful starters can include the reaction products between epichloprohydrin and the above mentioned starter compounds.

WO 93/24544 PCT/US93/04872

PParticularly preferred starter alcohols are primary and secondary 1 monohydric alcohols which are branched chain such as ethanol, proppanol, butanol, pentanol, hexanol, heptanol, octanol, nonanol, decanool, undecanol, dodecanol, tridecanol, tetradecanol, pentadecanol, hhexadecanol, octadecanol, isopropyl alcohol, 2-ethylhexanol, ssec-butanol, isobutanol, 2-pentanol, 3-pentanol and isodecanol. Parrticularly suitable alcohols are linear and branched primary alcohols (including mixtures) such as produced by the "Oxo" reaction oof C<sub>3</sub> to C<sub>20</sub> olefins. The alcohols may also be cycloaliphatic ssuch as cyclopentanol, cyclohexanol, cycloheptanol, cyclooctanol, as s well as aromatic substituted aliphatic alcohols such as benzyl alcohol, phenylethyl alcohol, and phenylpropyl alcohol. Other aliphatic c structures include 2-methoxyethanol and the like.

PPhenols include alkylphenyls of up to 30 carbons such as phenol, methhylphenol, ethylphenol, butylphenol, heptylphenol, octylphenol, noonylphenol, dinonylphenol and decylphenol. The aromatic radicals may contain other substituents such as halide atoms.

AAlcohols (polyols) having 2 or more hydroxyl groups, e.g., about two t to six hydroxyl groups and have 2 to 30 carbons, include glycols s such as ethylene glycol, propylene glycol, butylene glycol, pentylenne glycol, hexylene glycol, neopentylene glycol, decylene glycol, diethylene glycol, triethylene glycol and dipropylene glycol. Other poolyols include glycerine, 1,3-propanediol, pentaerythritol,l, galactitol, sorbitol, mannitol, erythritol, trimethylolethaane and trimethylolpropane.

TThe complex hydrophobe compounds can be prepared using a number of different methods. Three methods are described below, but synthhesis of these structures is not limited to just these methods.

TThe first method involves reaction of an unsubstituted or substituted alcohol or phenol, such as nonylphenol for example,

WO 93/24544 PCT/US93/04872

4417 30 --- 44

- 27 -

with an epoxide-e-containing hydrocarbon residue, such as nonylphenol glyocidyl ether. The reaction can be carried out under anhydrous condititions in the presence of at least 0.1 weight % but not more than 20 weight % of a suitable catalyst based on the starting weight of alcohold or phenol. Such a suitable catalyst can be, for example, potassisium hydroxide. Use of a suitable azeotroping solvent, such as toluene ( or cyclohexane, can aid in removal of water from the system. The reaction temperature should be at least about 60°C and at most the boiling point of the solvent, unless the reaction is done under pressure. . The reaction can be carried out under vacuum or in the absence of a s solvent. The catalyst and the alcohol can be added to the solvent, and I essentially all water can be removed, typically through azeotropping. Once the reaction temperature is reached, the epoxide can be addded. The product can be isolated by removal of solvent. An example of this type of procedure for producing a complex hydrophhobe is described in Example 1.

A second method involves reaction of a polymerizable cyclic ether, such as an epoxide, with a suitable polymerization catalyst like borron trifluoride in the presence of an initiating alcohol or phenol, referreed to as the starter. The reaction can be done in a suitable polymerrization solvent such as cyclohexane, which can be dried prior to pollymerization/oligomerization through its azeotrope. The reaction cann be carried out in the absence of a solvent. In a typical preparaticion, the catalyst, usually added at a level of from 0.1 mole % to 50 moble %, preferably from 0.1 mole % to 10 mole % based on the starting eppoxide, and the starter, usually added at a level of from 5.0 mole % to 100 mole %, preferably from 5.0 mole % to 50 mole %, based on the s starting epoxide, can be added to the solvent. The reaction temperature can be as low as -78°C to as high as the reflux temperature of tithe solvent at atmospheric pressure, but more preferably from 225°C to 75°C. The cyclic ether can then be added to the catalyst soluttion and polymerized. An example of this type of

Hage 30 of 115

procedure for proroducing a complex hydrophobe is described in Example 5.

At third method involves the reaction of an unsubstituted our substituted alcohol or phenol, such as nonylphenol or hexadecanol, , with a functional epoxide, such as epichlorohydrin. The starting alcohol or phenol can be added to a suitable reaction solvent, such as a toluene, along with potassium hydroxide, a tetraalkyl ammonium halide, such as tetraethylammonium iodide, and water. The a reaction can be carried out in the absence of a solvent. The reaction temperature can be regulated at about 65°C at which time the epichlorohydrin can be added. Varying the amount of epichlorohydrin and other ingredients can result in the formation of various dendriritic-like molecules. An example of this type of procedure for prooducing a complex hydrophobe is described in Example 8.

The alkoxylation is conducted using a catalytically-effective amount t of a catalyst, e.g., about 0.01 to 10, often about 0.5 to 5, weight percent based on the weight of a starter component, e.g., a complex hydrophhobe compound having at least one active hydrogen. The catalysts should retain their activities during the alkoxylation, regardless of thee amount of alkylene oxide employed. Thus, the amount of catalyyst can be based on the amount of starter provided to the alkoxylation 1 zone and not the degree of alkoxylation to be effected.

Illlustrative alkoxylation catalysts useful in this invention includde, for example, potassium hydroxide, diethylaluminum fluoride, oxides and hydroxides of barium and strontium, basic salts of alkaline earth metals especially calcium, calcium oxide (qquicklime), calcium hydroxide (slaked line), calcium sulfate, and the l like. Preferred catalysts for obtaining alkoxylation product mixturees having a narrow molecular weight distribution are disclosed in U.S.3. Patent 4,754,075, the disclosure of which is

incorporated herein by reference. This invention is not intended to be limited in any mmanner by the permissible catalysts.

Normally, the catalyst and the starter component are admixed and theen the alkylene oxide is added at the reaction temperature until the desired amount of alkylene oxide has been added, then the product is neutralized and can be finished, if desired, in any procedure including stripping unreacted starter material from the productt mixture, filtration, or further reaction.

The temperature of the alkoxylation is sufficient to provide a suitablile rate of reaction and without degradation of the reactants or reaction products. Often, the temperatures range from between about 500°C and 270°C, e.g. from about 100°C to 200°C. The pressure may alsso vary widely, but when low-boiling alkylene oxides such as ethylenee oxide and propylene oxide are employed, a pressurized reacctor is preferably used.

The alkoxylation reaction medium is preferably agitated to ensure a good dispersal of the reactants and catalyst throughout the reaction medium. Also, the alkylene oxide is usually added at a rate approximating that which it can be reacted.

Neutralization may assist in the recovery of the catalyst from thee alkoxylation product mixture. When neutralizing, acids that may teend to form catalyst-containing gel structures or solids that clog fifiltering apparatus should be avoided. Conveniently, sulfuric acid, phoosphoric acid, propionic acid, benzoic acid and the like are used.

Th'he above procedure represents but one of many equally viable veersions of suitable alkoxylation processes. Other versions are posssible through combinations of the options available in the various proceess steps.

Thhe catalytic alkoxylation reactions of this invention can be effected, fifor example, by conventional methods such as (1) batch processes; (2) continuous fixed-bed processes; and (3)

WO 93/24544 PCT/US93/04872

- 30 -

continuous fluidized reactor processes. In a batch reactor, the catalyst is kept t suspended in the reactant by shaking or stirring. In a fluidized reacctor, the catalyst is at a particular original level. As the velocity of tithe reactant stream is increased, the catalyst bed expands upward to a second level, and at a critical velocity it enters into violent turbulence. The fluidized reactor is particularly useful for removing orr supplying the heat necessary to maintain a fixed catalyst temperature. The fluidized reactor can usually be employed only on a ratheer large scale since good fluidization requires a reactor larger than about 1.5 inches in diameter.

AAs indicated above, the end groups described by  $R_6$ ,  $R_{11}$ ,  $R_{14}$  and  $RR_{20}$  can be a variety of structures including sulfates. The sulfation obf a hydroxyl containing end group can be carried out using one of seeveral methods, such as with  $SO_3$  or sulfamic acid.

In the sulfation reaction using sulfamic acid, a complex hydropphobe alkoxylate, for example ethoxylated 1,3-bis(nonylphenoxy)-2-propanol, is dried at high temperature, at or about 100°C forr example, and low pressure, at or below 50 mm Hg for example. The ethoxylate is cooled to a reaction temperature at or about 60°C, forr example, and sulfamic acid is charged. The reaction temperature is kept at or about 110°C. Once the reaction is complete, an amount of an mono- or disubstituted alkanolamine can be added to remove residual acid, and prevent hydrolysis upon introduction of water. A dilutiting solvent, such as ethanol and/or water, can be added along with the alkanolamine. An example of a preparation of a sulfate using sulfamic acid is shown in Example 11.

The alkoxylation products produced by the alkoxylation processes of this invention, including the complex hydrophobe compounds having at least one active hydrogen, can undergo further reaction(s) to afford desired derivatives thereof. Such permissible derivatization 1 reactions can be carried out in accordance with conventional procedures known in the art. Illustrative derivatization

WO 93/24544 PCT/US93/04872

- 31 -

reactions includde, for example, esterification, etherification, amination, alkyllation, hydrogenation, dehydrogenation, reduction, acylation, condeensation, carboxylation, oxidation, silylation and the like, including ppermissible combinations thereof. This invention is not intended to t be limited in any manner by the permissible derivatization recactions or permissible derivatives of alkoxylation products and coomplex hydrophobe compounds.

MMore particularly, the hydroxyl-terminated alkoxylation prooducts and complex hydrophobe compounds of this invention can unndergo any of the known reactions of hydroxyl groups illustrative of which are reactions with acyl halides to form esters; with ammonia, a nitrile, or hydrogen cyanide to form amines; with alkyl acid sulfattes to form disulfates; with carboxylic acids and acid anhydrides to foorm esters and polyesters; with alkali metals to form salts; with ketennes to form esters; with acid anhydrides to form carboxylic acids;; ring-opening reactions with lactones, tetrahydrofuran; dehydrogenationn to form aldehydes, isocyanates to form urethanes, and the like.

Alkkoxylation product mixtures prepared by the processes of this invention comprise alkoxylation species that can be represented by tithe formulae (i) and/or (ii) above. The alkoxylation product mixturees of this invention can be characterized as the condensation reaction products of alkylene oxides and complex hydrophobe compounds having at least one active hydrogen in which the mole ratio off reacted alkylene oxide per active hydrogen is at least about 4, say, about 4 to 16 or 24, preferably about 5 to 12. The product mixtures can have at least one alkoxylation moiety which constitutes at least about 10,0, say, about 20 to 30 or 40, and most often about 20 to 30, weight perceent of the composition. The alkoxylation mixtures of this invention allso can have a relatively symmetrical distribution. Hence, the porticion of the product mixture having three or more

oxyalkylene uniit groups (per active hydrogen site of the complex hydrophobe compound) than the peak alkoxylation specie is relatively minor, e.g., ofteen less than about 12, say, less than 10, and often about 1 to 10, wweight percent of the mixture. Similarly, the alkoxylation species having fewer oxyalkylene groups (per active hydrogen site off the complex hydrophobe compound) by three or more oxyalkylene groups from the average peak alkoxylation species is usually relatively minor, e.g., less than about 15, say, less than about 10, often about 60.5 to 10, weight percent of the composition. Generally, the cone oxyalkylene unit higher and the one oxyalkylene unit lower alkoxylates in respect to the most prevalent alkoxylation specie are preseent in a weight ratio to the most prevalent alkoxylation specie of about 00.6:1 to 1:1.

The nonionic surfactant compounds encompassed by formulae (i) andd (I) may be converted to anionic surfactant compounds by chemically modifying a terminal or pendant hydroxyl group thereof. (One such approach includes the steps of reacting the nonionic surfactant compounds encompassed by formulae (i) and (I) with sulfuric accid or sulfamic acid and then optionally converting the resulting free sculfate esters (when sulfuric acid is used) or ammonium saltits (when sulfamic acid is used) into corresponding alkali metal or r alkanolamine salts such as monoethanolamine by conventional porocedures.

AAnother approach includes the steps of reacting the nonionic surfactant compounds encompassed by formulae (i) and (I) with phosphoruus pentoxide or phosphorus oxychloride and then optionally conveerting the resulting free phosphate esters into corresponding scalts with alkali metal, ammonium or alkanolamine salts whereby mmixtures of mono-esters and di-esters are obtained. These mixturess may be used as an anionic surfactant without being isolated into theeir constituents.

A further approach includes the steps of reacting the nonionic surfact tant compounds encompassed by formulae (i) and (I) with maleic anhaydride, reacting resultant maleate esters with an alkali metal bisuulfite and then optionally converting the resulting alkali metal sulfifosuccinates into corresponding free acids or ammonium or alilkanolamine salts. When at least one mole of maleic anhydride is useed relative one mole of the compounds of the formula (i) and (I) in the e above reaction, mixtures of half ester isomers having a sulfo group attached to either alpha or beta position relative to the terminal carboxyyl group. These mixtures may also be used as an anionic surfactant without isolation. Conversely, when at least two moles of the compounds of the formulae (i) and (I) are reacted with one mole of maldeic anhydride, di-esters are obtained.

Preferred macromonomer compounds useful in this invention includde those represented by the formulae:

$$R^{1}$$
— $(OR^{2})z$ — $OC(O)$ — $NH$ — $C(CH_{3})_{2}$ — $C(CH_{3})$ = $CH_{2}$ 
(XII)

$$\begin{array}{c} R^4 \\ | \\ R^1\text{-}(OR^2)_z\text{-}CH_2OCHCH_2OCH_2C=CH_2 \\ \\ | | \\ (((OR_{19})_jOH) \end{array} \tag{XIII}$$

- 34 -

 $R^4$ 

 $R^{1}$  -  $(OR^{2})_{z}$  - OCC(O)-C= $CH_{2}$ 

(XIV)

wherein R<sup>1</sup>, R<sup>2</sup>, R<sup>4</sup>, R<sub>19</sub>, z and j are as defined herein.

The macromonomer compounds useful in this invention can undergo fuurther reaction(s) to afford desired derivatives thereof. Such permissibble derivatization reactions can be carried out in accordance with conventional procedures known in the art.

Illustrative derrivatization reactions include, for example, esterification, eetherification, alkoxylation, amination, alkylation, hydrogenation, dehydrogenation, reduction, acylation, condensation, carboxylation, coxidation, silylation and the like, including permissible combinations thereof. In an embodiment of this invention, the macromonomers can contain carboxyl or other ionic or acid moieties. This invention is not intended to be limited in any manner by the permissible derivatization reactions or permissible derivatives of macromonomer compounds.

Mdore particularly, the hydroxyl-terminated macromonomers compounds of this invention can undergo any of the known reactions of hydroxyl groups illustrative of which are reactions with a acyl halides to form esters; with ammonia, a nitrile, or hydrogen cyanide to form amines; with alkyl acid sulfates to form disulfates; with carboxylic acids and acid anhydrides to form esters and polyesters; with alkali metals to form salts; with ketenes to form esters; with acid anhydrides to form carboxylic acids; with oxygen to form aldehydess and carboxylic acids; ring-opening reactions with lactones, tetrahhydrofuran; dehydrogenation to form aldehydes, isocyanates to form urethanes, and the like.

The monoethylenically unsaturated macromonomer component is scubject to considerably variation within the formula presented previously. The essence of the maromonomer is a complex

- 35 -

hydrophobe carrrying a polyethoxylate chain (which may include some polypropoxxylate groups) and which is terminated with at least one hydroxy group. When the hydroxy-terminated polyethoxylate complex hydrophhobe used herein is reacted with a monoethylenically unsaturated modnoisocyanate, for example, the result is a monoethylenicallyly unsaturated urethane in which a complex hydrophobe polyyethoxylate structure is associated with a copolymerizable a monoethylenic group via a urethane linkage.

Thee monoethylenically unsaturated compound used to provide the monnoethylenically unsaturated macromonomer is subject to wide vvariation. Any copolymerizable unsaturation may be employed, such as acrylate and methacrylate unsaturation. One may also use allilylic unsaturation, as provided by allyl alcohol. These, preferablyly in the form of a hydroxy-functional derivative, as is obtained by reaccting a C2-C4 monoepoxide, like ethylene oxide, propylene oxide or butylene oxide, with acrylic or methacrylic acid to form an hydroxyy ester, are reacted in equimolar proportions with an organic compounnd, such as toluene diisocyanate or isophorone diisocyanate. The preferred monoethylenic monoisocyanate is styryl, as in alpha, alpha-dimethyl-m-isopropenyl benzyl isocyanate. Other suitable organice compounds include, for example, monoethylenically unsaturated estiters, ethers, amides, ureas, anhydrides, other urethanes and t the like.

Thee polymers of this invention can be prepared via a variety of polymnerization techniques known to those skilled in the art. The technique of polymerization influences the microstructure, monomer sequence distribution in the polymer backbone and its molecular weight to influence the performance of the polymer. Illustrative polymerization techniques include, for example, conventional annd staged emulsion polymerization via batch, semicontinuous, or a continuous processes, micellar polymerization, inverse emulsion polymerization, solution polymerization, non-

aqueous dispersion polymerization, interfacial polymerization, emulsion polymerization, suspension polymerization, precipitation polymerizationn, addition polymerizations such as free radical, anionic, cationnic or metal coordination methods, and the like.

Th'he thickeners of this invention possess structural attributes of twwo entirely different types of thickeners (those which thicken by pH ( dependent solubilization, e.g., alkali solubilization, of a high molecular weight entity, and those which thicken due to association), annd this may account for the superior thickener properties which are obtained herein.

The aqueous emulsion copolymerization is entirely conventional. To obtain an estimate of thickening efficiency, the product can be a diluted with water to about 1% solids content and then adjust the pH t to solubilize the polymer or neutralized with alkali. The usual alkaali is ammonium hydroxide, but sodium and potassium hydroxide, and even amines, like triethylamine, may be used for neutralization. The usual acid is sulfuric acid, acetic acid and the like. After neutralization or adjustment to appropriate pH, the product dissolves in the water to provide an increase in the viscosity. In the normal mode of addition, the unsolubilized or unneutralized t thickener is added to a paint and then the pH is adjusted or neutralized. This facilitates handling the thickener because it has a lower viscosity before pH adjustment or neutralization... This procedure also makes more water available for the paint formulation.

This polymers of this invention are preferably produced by conventionaal aqueous emulsion polymerization techniques, using appropriate emulsifiers for emulsifying the monomers and for maintaining the polymer obtained in a suitable, dispersed condition. Commonly useed anionic surfactants such as sodium lauryl sulfate, dodecylbenzenee sulfonate and ethoxylated fatty alcohol sulfate can be

- 37 -

used as emulsififiers. The emulsifier may be used in a proportion of 1/2 to 6% of the a weight monomers.

Preeferably, water-soluble initiators such as alkali metal or ammonium ppersulfate are used in amounts from 0.01 to 1.0% on the weight of mnonomers. A gradual addition thermal process employed at temperatures between 60°C to 100°C is preferred over redox systems.

Thee polymerization system may contain small amounts (0.01 to 5% by wweight, based on monomer weight) of the chain transfer agent remercaptans such as hydroxyethyl mercaptan, 6-mercaptopropiopnic acid and alkyl mercaptans containing from about 4 to 22 carbon automs, e.g., ethyl hexyl mercapto propionate and tertiary dodecyl mercaptan, and the like. The use of mercaptan modifier reduces the molecular weight of the polymer and therefore its thickening exefficiency.

Thee polymers of this invention may further be modified by introducing an amount of component (D), namely, one or more polyethylenicallily unsaturated copolymerizable monomers effective for crosslinkings, such as diallylphthalate, divinylbenzene, allyl methacrylate, thrimethylol propane triacrylate, ethyleneglycol diacrylate or dimethacrylate, 1,6-hexanediol diacrylate or dimethylacrylatite, diallyl benzene, and the like. Thus, from about 0.05 or less to abbout 20% or greater of such polyethylenically unsaturated compound based on total weight of monomer may be included in the ecomposition forming the polymer. The resulting polymers are eitither highly branched or in the form of three-dimensional nettworks. In the neutralized salt form, those networks swell in an aquaeous system to act as a highly efficient thickener.

Other illustrative polyethylenically unsaturated monomers useful in this invention include, for example, any copolymerizablee compound which contains two or more nonconjugated points of ethylenic unsaturation or two or more

- 38 -

nonconjugated vinylidene groups of the structure, CH2=C=, such as divinyltoluene, , trivinylbenzene, divinylnaphthalene, trimethylene glycol diacrylatte or dimethacrylate, 2-ethylhexane-1,3-dimethyacrylatte, divinylxylene, divinylethylbenzene, divinyl ether, divinyl sulfone,; allyl ethers of polyhdric compounds such as of glycerol, pentaeerythritol, sorbitol, sucrose and resorcinol, divinylketone, c divinylsulfide, allyl acrylate, diallyl maleate, diallyl fumarate, diallyl phthalate, diallyl succinate, diallyl carbonate, diallyl malonatte, diallyl oxalate, diallyl adipate, diallyl sebacate, diallyl tartrate,; diallyl silicate, triallyl tricarballylate, triallyl aconitate, triallyl citrate, triallyl phosphate, N,N-methylenediaccrylamide, N,N'-methylenedimethacrylamide, N,N'-ethylidenediaccrylamide and 1,2-di-(a-methylmethylenesulfonamide)-ethylene.

Thhe polymer may be utilized in a variety of ways to provide the thickener or thickened compositions of this invention. For example, the polymer, while in aqueous dispersion or dry form, may be blendedd into an aqueous system to be thickened followed by addition of a pbH adjusting agent, e.g., neutralizing agent. Alternatively, t the polymer may first be solubilized or neutralized in aqueous disperssion form and then blended with the aqueous system. Preferably, if cco-thickening by a surfactant is desired, the components aree separately blended (as dry components or as dispersions or & slurries) into an aqueous dispersion to be thickened, followed by thee pH adjustment step, e.g., neutralization step. Although aqueeous concentrates of the polymer in unsolubilized form, e.g., in acid forrm, and the surfactant may be formed and added to an aqueous disperrsion to be thickened as needed, followed by pH adjustment, e.gg., neutralization, such concentrates tend to be too viscous for easyy handling. It is nevertheless possible to prepare either a dry bldend or an aqueous, high solids composition which is sufficiently loww in viscosity as to be pumpable or pourable, and then to

Page 41 of 11!

further thickeen the admixture by addition of alkaline or acidic materials.

The polymer thickener may be provided in a dry state in number of wayys. For example, the unsolubilized polymer may be spray or druma dried and, if desired, blended with a surfactant cothickener. However, it is also possible to spray dry or otherwise dehydrate the solubilized polymer thickener, and then reconstitute the aqueous thhickener dispersion at a future time and place by agitation in a saqueous medium, provided the pH of the dispersion is maintained at t an appropriate value, e.g., at pH 7 or higher for neutralizations.

of this invention for aqueous thickening is to add the aqueous dispersion of the polymer to the medium to be thickened and, after mixing, to introduce an alkaline or acidic material to ionize component (A).). The major portion of the thickening effect is usually obtained in a fifew minutes upon pH adjustment. In the presence of high concentrations of electrolytes, the viscosity development may take much longger. This method of applying a polymer to an aqueous system before I pH adjustment enables one to handle a high solids thickener in a 1 non-viscous state, to obtain uniform blend, and then to convert to a highly viscous condition by the simple addition of an alkaline or aciddic material to bring the pH of the system to an appropriate value.

Th'he aqueous solutions thickened with the neutralized polymers of this invention exhibit good viscosity stability even at a pH as high as 13. The polymer may be used to thicken compositions under acidic coonditions in the presence of a relatively large amount of surfactants wherein the thickened composition, for example, an aqueous system, has a pH below 7, even as low as 1.

Ann enhancement of thickening (herein termed "cothickening") can result upon the addition of a surfactant to an

- 40 -

aqueous systemm containing the polymer of this invention, when the polymer is solubilized. In some cases the thickening can be enhanced up to 0 about 40 times the viscosity afforded by the solubilized polymer alone. A wide range of surfactants may be used. Although trace amounts ( of surfactant may be residually present from the polymerization 1 of the monomers comprising the polymer (for example, whateever may remain of the about 1.5 weight percent surfactant on nmonomers), such amounts of surfactant are not believed to resuult in any measurable co-thickening.

Onn the basis of an aqueous system containing about 0.1 to 5% by weight of polymer solids, a useful amount of surfactant for optimum co-thickening is about 0.1 to 1.0% by weight of the total system. As inddicated, the amounts of polymer and surfactant cothickener many vary widely, even outside these ranges, depending on polymer and surfactant type and other components of the aqueous system to be thickened. However, the co-thickening can reach a maximum as scurfactant is added and then decreases as more surfactant is added. Hence, it may be uneconomical to employ surfactant in amounts outside the stated concentrations and polymer/surfactant ratios, but this can be determined in a routine manner in each case.

The preferred method of application of the polymer and the surfactant f for aqueous thickening is to add in any sequence the polymer and the surfactant to the medium to be thickened and, after mixing, to introduce an alkaline or acidic material to adjust the pH. This method of f applying polymer and surfactant to an aqueous system before ppH adjustment enables one to handle a high solids thickener in a r non-viscous state, to obtain a uniform blend, and then to convert to a l highly viscous condition by the simple addition of an alkaline or aciddic material to bring the pH of the system to an appropriate value. However, the pH of the aqueous system may also be adjusted beftfore addition of the surfactant.

Page 43 01 11

The e surfactants which may be used include nonionics and anionics, siningly or in combination, the selection necessarily depending upon a compatibility with other ingredients of the thickened or thickenable dilispersions of this invention. Cationic and amphoteric surfactants may also be used provided they are compatible with a the polymer and other ingredients of the aqueous system, or are used in such small amounts as not to cause incompatibility.

Suititable anionic surfactants that may be used include the higher fatty r alcohol sulfates such as the sodium or potassium salt of the sulfates of f alcohols having from 8 to 18 carbon atoms, alkali metal salts or ammine salts of high fatty acid having 8 to 18 carbon atoms, and sulfo onated alkyl aryl compounds such as sodium dodecyl benzene sulfonatte. Examples of nonionic surfactants include alkylphenoxypolylyethoxyethanols having alkyl groups of about 7 to 18 carbon atoms and about 9 to 40 or more oxyethylene units such as octylphenoxy-polylyethoxyethanols, dodecylphenoxypolyethoxyethanols; ethylenne oxide derivatives of long-chain carboxylic acids, such as lauric, nmyristic, palmitic, oleic; ethylene oxide condensates of long-chain alcohols such as lauryl or cetyl alcohol, and the like.

Examples of cationic surfactants include lauryl pyridinium chloride, octylbenzyltrimethyl-ammonium chloride, dodecyltrimethyl-lammonium chloride condensates of primary fatty amines and ethylylene oxide, and the like.

The e foregoing and numerous other useful nonionic, anionic, cationic,c, and amphoteric surfactants are described in the literature, such as McCutcheon's Detergents & Emulsifiers 1981 Annual, North AAmerica Edition, MC Publishing Company, Glen Rock, NJ 074522, U.S.A., incorporated herein by reference.

In ggeneral, solvents and non-solvents (or mixtures of solvents, non-solvents other organics and volatiles) can be used to manipulate the vviscosity of polymer containing systems. In the

examples hereisin, it is interesting to note how mineral spirits act like co-thickener, annd how the water solubility of the other solvent influences how v much mineral spirits can be added before the solution separates into a two phase system. The co-thickening with mineral spirits s has utility in textile printing pastes, and in waterborne autitomotive basecoats. These systems usually contain mineral spirits s (because of the pigments used therein), so that the mineral spirits s provide an economical way of increasing viscosity and improving the efficiency of the thickener.

The amount of the polymer that may be dissolved in any given aqueous composition may fall within a wide range depending on the particular viscosity desired.

Thhus, although any effective amount of the polymer may be employed forr dissolution, typically from about 0.05 to about 20%, preferably from about 0.1 to about 5%, and most preferably from about 0.1 to about 3% by weight, based on the weight of the final aqueous composition inacluding polymer is used.

Foor latex paint compositions, the polymer may be dissolved thereisin in an amount of from about 0.05 to about 5%, and preferably from about 0.1 to about 3% by weight, based on the weight of the total composition including polymer.

Thhe polymers of this invention may be employed as thickeners for controlling viscosity of any aqueous based composition. An aqueous based composition is an aqueous composition as herein defined to be a composition wherein water comprises at least 10% by weight of the total composition (including 100% water).

Foor example, aqueous dispersions, emulsions, suspensions, solutions, slurries and the like, may be thickened by the polymers of this invention.

Tyypical aqueous compositions include compositions to be applied to textililes such as latex adhesives, warp sizes, backings for rugs and other r pile fabrics. The polymer may also be used when

7 6Ac 46 6 1 1 1 1

・とし シン・ニオンササ

thickening is dessired in the purification of raw water such as the saline water useed in the recovery of oil from exhausted oil wells by water flooding teechniques. Other aqueous coatings compositions to which the polymmer can be added for thickening purposes include drilling muds, exaulks, adhesives, coating compositions such as paper coatings, I furniture finishes, ink compositions, latex paints, foundary core weashes, and the like.

Preferably, the polymer is used to thicken aqueous coating compositions, and more preferably latex paint compositions. Examples of suititable latex paint compositions include those based on resins or binders of acrylonitrile, copolymers of acrylonitrile wherein the comonomer i is a diene like isoprene, butadiene or chloroprene, homopolymers otof styrene, homopolymers and copolymers of vinyl halide resins such as vinyl chloride, vinylidene chloride or vinyl esters such as vivinyl acetate, vinyl acetate homopolymers and copolymers, copoolymers of styrene and unsaturated acid anydrides like maleic anhydrides, homopolymers and copolymers of acrylic and methacrylic acidd and their esters and derivatives, polybutadiene, polyisoprene, buttyl rubber, natural rubber, ethylene-propylene copolymers, olefifins resins like polyethylene and polypropylene, polyvinyl alcohobl, carboxylated natural and synthetic latices, epoxies, epoxy esters andd similar polymeric latex materials.

Lattex paint compositions are well known in the art and typically comprise an emulsion, dispersion or suspension of discrete particles of resinn binder and pigment in water. Optional ingedients typically include a thickeners, antifoam agents, plasticizers, surfactants, coallescing agents, and the like.

Thee polymers described herein are useful in a variety of aqueous systemss, such as textile coatings (woven and nonwoven), latex paint formulations, cosmetic formulations, pigment dispersions and I slurries, dentrifrices, hand lotions, liquid detergents, quenchants, agricultural chemicals, concrete additives,

transmission flduids, waste water treatment (flocculants), turbulent drag reduction,, aircraft anti-icing, automation coatings (OEM and refinish), archititectural coatings, industrial coatings and the like.

Othther applications include, for example, paper making, mineral processing, brine viscosification, superabsorbency, enhanced oil recovery, personal care products, biomedical, pharmaceuticall and the like.

The macromonomers of this invention are particularly useful as an emmulsifier in the emulsion or dispersion polymerization of ethylenicallyy unsaturated monomers. Examples of monomers include acrylic: monomers such as acrylic acid, methyl acrylate, butyl acrylate, 2 ethylhexyl acrylate, methyl methacrylate, acrylonitrile, accrylamide and 2-hydroxethyl acrylate; aromatic monomers such as styrene and divinylbenzene; vinyl esters such as vinyl acetate; hhalogen-containing monomers such as vinyl chloride and vinylidene; chloride; conjugated diene monomers such as butadiene, isopprene and chloroprene; ethylene, maleic anhydride and methyl mahleate.

AAny known initiator such as hydrogen peroxide, potassium persulfate, azobisisobutyronitrile and benzoyl peroxide may be used in 1 the emulsion polymerization in combination with a known polymerization promoter such as sodium bisulfite and ammonium ferrous sulfate.

TThe macromonomers of this invention may be used in an amount of 0.1.1 to 20%, preferably 0.2 to 5.0% by weight of the total monomers singly or in combination with a conventional emulsifiers or protective coblloid.

TThe resulting polymer emulsions or dispersion may be applied to woodds, metals, paper, textiles, and concrete structures as an adhesive, copating agent or reinforcing agent.

TThe macromonomer compounds of this invention find use not only as s an emulsifier or dispersant of various materials such

as monomers in 1 the emulsion or suspension polymerization, waxes, dyes, pigments and pesticides, but also as latex binders and as post-treating agents f for textile materials and antistatic agents for plastic products. By thee use of the macromonomers of this invention, defects of conventional s surfactants remaining after use may be alleviated.

The e macromonomer compounds of this invention can be homopolymerized or copolymerized with conventional monomers via a variety of polymerization techniques known to those skilled in the art. The technique of polymerization influences the microstructure, monomer sequence distribution in the polymer backbone and its molecular weight to influence the performance of the polymer. Illustrative polymerization techniques include, for example, conventional and staged emulsion polymerization via batch, semicontinuous, or continuous processes, inverse emulsion polymerization, solution polymerization, non-aqueous dispersion polymerization, interfacial polymerization, emulsion polymerization, suspension polymerization, precipitation polymerization, addition polymerizations such as free radical, anionic, cationic or metal coordination metahods, and the like.

The surfactant compounds of this invention can be used wherever tithere is a need to increase surface activity. They are particularly valuable where process conditions demand the greatest efficiency from vwater, e.g., in wetting, penetrating, emulsifying, lubricating and a dispersing. Illustrative applications include, for example, agriculature, e.g., emulsifiers and wetters, herbicidal adjuvants and fifruit washes; cleaners, e.g., household detergents, dry cleaning detergents, industrial hand cleaners, detergent/sanitizers, solvent degreaseers, metal cleaners, hard surface cleaners, commercial launndry detergents and car wash detergents; dust wetting, e.g., copal mines, ceramic industries and foundries; leather, e.g., hidesoakings, degreasing, fatliquor stabilization, tanning and dyeing; paint, e.g., emulsion polymerization of latexes, latex

- 46 -

stabilization, pigigment wetting and dispersion; paper, e.g., pulping, absorbent papeers, de-inking waste water, wet-felt washing and adhesives; and I textiles, e.g., greige goods scouring, bleaching, carbonizing of wwool, scouring of cotton, wool and polyester/cotton blends, wetting g and rewetting agents, desizing of cotton and polyester/cotton blends, polyethylene softener emulsifiers, mineral oil and solvent emulsifiers, fiber lubricant emulsifiers, emulsion polymerization, i, dyeing and finishing of synthetics, naturals and blends, dye leveling agents and resin bath additives.

Addditionally, the compounds of this invention may be useful as compoonents of sanitizing solutions for use on food processing equipipment and utensils and on beverage containers, including milk containers and equipment; emulsifiers and/or surface active agents in the manufacture of articles or components of articles intendeed for use in producing, manufacturing, packing, processing, prepparing, treating, packaging, transporting or holding food; inert (or occasionally active) ingredients for use in formulations applied to growwing crops or to raw agricultural commodities after harvest; and ineert (or occassionally active) ingredients for use in formulations applied to animals.

As 3 used herein, the term "complex hydrophobe" is contemplated too include all permissible hydrocarbon compounds having 2 or monre hydrophobe groups, e.g., bis-dodecylphenyl, bisnonylphenyl, bisis-octylphenyl and the like.

Forr purposes of this invention, the term "hydrocarbon" is contemplatedd to include all permissible compounds having at least one hydrogen annd one carbon atom. In a broad aspect, the permissible hyddrocarbons include acyclic and cyclic, branched and unbranched, caarbocyclic and heterocyclic, aromatic and nonaromatic organic compounds which can be substituted or unsubstituted.

- 47 -

As uused herein, the term "substituted" is contemplated to include all peermissible substituents of organic compounds unless otherwise indicatted. In a broad aspect, the permissible substituents include acyclic aand cyclic, branched and unbranched, carbocyclic and heterocyclic,:, aromatic and nonaromatic substituents of organic compounds. Illustrative substituents include, for example, alkyl, alkyloxy, aryl, anryloxy, hydroxy, hydroxyalkyl, amino, aminoalkyl, halogen and the elike in which the number of carbons can range from 1 to about 20 or nmore, preferably from 1 to about 12. The permissible substituents can a be one or more and the same or different for appropriate organic compounds. This invention is not intended to be limited in any mnanner by the permissible substituents of organic compounds.

The e invention is illustrated by certain of the following examples.

## Example 1

## Preparation of 1.3-Bis(nonylphenoxy)-2-propanol

To aa five neck, two liter round bottom flask equipped with an addition funnel, thermometer, nitrogen dispersant tube, mechanical stirreer, and a decanting head with a water-cooled condenser were aadded 220 grams (1.00 mole) of nonylphenol and 250 milliliters of cycldohexane. The solution was then heated to reflux and 2.8 grams (11.3 wt. % based on nonylphenol) of potassium hydroxide in 10 nmilliliters of water was slowly added to the flask. After essentially y all the water was recovered in the decanting head (10 milliliters + 11 milliliter formed), 250.7 grams (0.91 mole) of nonylphenyl glyccidyl ether as added dropwise. During the addition of the glycidyl etheer, the reaction temperature was maintained between 60 and 80°C. Aftiter the addition was complete, the solution was refluxed for four r hours. The contents of the flask were then washed with a five perceent aqueous solution of phosphoric acid, and the

organic layer was separated from the water layer and washed twice with deionized water. The reaction mixture was then placed in a one liter round bottdom flask, and the remaining cyclohexane and unreacted nonyl/lphenol were recovered by distillation, first at atmospheric preessure, then under vacuum at 0.2 mm Hg. The kettle temperature was not allowed to exceed 180°C during the distillation to prevent discobloration of the product. The concentrated solution was then refiltered to give 425 grams of a pale-yellow liquid. Endgroup MW analylysis gave a molecular weight of 506.8 (theoretical MW = 496.8). Ir andd nmr spectra were identical to previously recorded spectra for the compound.

## Example 2

## Preparation of 1.3-Bis(nonylphenoxy)-2-propanol

To a five neck, two liter round bottom flask, equipped with an addition funnel, thermometer, nitrogen dispersant tube, mechanical stirrrer, and a decanting head with a water-cooled condenser, weree added 300 milliliters of cyclohexane and 451.7 grams (2.05 mole) of naonylphenol. The solution was then heated to reflux and 58.9 grams s (1.05 mole) of potassium hydroxide in 60 milliliters of water was slowlyly added via the addition funnel. After essentially all the water was recovered in the decanting head (60 milliliter + 19 milliliters formaed), the reaction was cooled to 40°C, and 92.5 grams (1.00 mole) of eppichlorohydrin was slowly added. During the addition, the resection temperature was maintained below 60°C by controlling the 1 rate of epichlorohydrin addition. After all the epichlorohydrinn was added, the solution was allowed to stir for one hour, and then i brought to reflux for an additional three hours. The reaction mixturre was then filtered under vacuum through a steamjacketed Buchnner funnel to remove the potassium chloride formed as a by-product. The filtration process was performed a total of three times to removee the majority of the salts. The reaction mixture was

- 49 -

then placed in a cone liter round bottom flask, and the remaining cyclohexane and curreacted nonylphenol were recovered by distillation, first at atmospheric pressure, then under vacuum at 0.2 mm Hg. The kettitle temperature was not allowed to exceed 180°C during the distillation to prevent discoloration of the product. The concentrated soluution was then refiltered to give 275 grams of a pale-yellow liquid. Erind-group MW analysis gave a molecular weight of 459.7 (theoretical MW = 496.8). Ir and nmr spectra were identical to previously recorded spectra for the compound.

## Example 3 Preparation of 5 Mole Ethoxylate of 1,3Bis(nonvlphenoxy)-2-propanol

To aa 500 milliliter, stainless steel, high pressure autoclave was chaarged 200 grams (0.40 mole) of 1,3-bis(nonylphenoxyy)-2-propanol, which contained a catalytic amount of the potassium sallt of the alcohol as described in Example 1. After purging the reacttor with nitrogen, the alcohol was heated to 130°C with stirring, and 86.9 grams (2.0 mole) of ethylene oxide was added over a two hour period. The reaction temperature and pressure were maintained from 1 130°C to 140°C and 60 psig during the course of the reaction. After the addition of ethylene oxide was complete, the reaction mixture was held at 140°C for an additional hour to allow all the ethylene oxidde to cook out. The reaction mixture was dumped while hot, under r nitrogen, and neutralized with acetic acid to yield 285 grams of a ppale-yellow liquid.

- 50 -

# Example 4 Preparation of Adduct of Nonviphenvi Glycidyl Ether and 5 Mole Ethoxylate of 1.3-Bis(nonylphenoxy)-2-propanol

To a five neck, one liter, round bottom flask equipped as in Example 1 was added 119.8 grams (0.17 mole) of the 5 mole ethoxylate of 1,33-bis(nonylphenoxy)-2-propanol and 100 milliliters of cyclohexane. The mixture was refluxed (100°C) for one hour to remove residualil water, and then cooled to 50°C under nitrogen to add 0.5 grams of BFF3/Et2O. Nonylphenyl glycidyl ether (46.0 grams, 0.17 mole) was then a added to the flask over a one hour period, and the reaction was hexated to reflux. After three hours at reflux, the reaction mixturge was transferred to a separatory funnel, while hot, and washed with a saturated aqueous solution of sodium bicarbonate. Th'he organic layer was separated from the water layer, and washed twirice with hot deionized water. The washes were performed at 500°C to facilitate the separation of the two layers. The water and cyclobhexane were then evaporated from the organic layer, under vacuum, to yield 145 grams of a pale-yellow, viscous liquid. End-group mole ecular weight analysis gave a molecular weight of 880 (theoretical mol·lecular weight = 993).

## Example 5

## Prepparation of Poly(nonylphenol glycidyl ether)

To a 500 milliliter round bottom equipped with an overhead stirreer, nitrogen inlet, reflux condenser, additional funnel, and temperature controller was charged 1.9 grams of ethanol (22 mmoles) and 2000 grams of cyclohexane. The solution was brought to 50°C. Once heaated, 0.5 milliliters (4 mmoles) of BF3/Et2O was added using a 2 millililiter syringe. Once the acid was added, 100.0 grams of nonylphenol glyycidyl ether (362 mmoles) was added dropwise so as to

- 51 -

maintain a reaction temperature of 45°C-55°C. Once the glycidyl ether was added, , the solution is refluxed for 3 hours, then cooled to about 50°C.

Whilile hot (<60°C) the organic was transferred to a separatory funnell and was washed once with 100 milliliters of 5% sodium bicarbonaate solution. The aqueous layer was drained and the organic was washhed two more times with 100 milliliter portions of deionized water. The aqueous layers were decanted and the organic was dried for at kleast 1 hour over magnesium sulfate. Once dry the magnesium sulfate was filtered from the organic which was stripped of solvent using aa rotary evaporator. The final yield of viscous polymer was 100) grams. The GPC molecular weight was Mw = 2600 and the Mn = 17000 based on monodisperse polystyrene standards.

## Example 6

## Ethoxyvlation of Poly(nonvlphenol glycidyl ether)

To a a 500 milliliter stainless steel Zipperclave was added 60.0 grams (0.0355 moles based on an approximate molecular weight of 1700 gram/mol·le) of the resin prepared in Example 5 along with 0.5 grams of potassiuum hydroxide. The vessel was attached to an automated ethoxyylation unit and was heated to 50°C. The vessel was continuously purgged with nitrogen for 15 minutes and was then heated to 100°C vwhere it was again continuously purged with nitrogen for anothther 15 minutes. The vessel was then heated to 140°C and was given a series of 6 purges by pressuring the vessel up to 80 psi, and then vennting. Once the venting process was complete, the vessel was pressuured to 20 psi with nitrogen.

The ethylene oxide lines were opened to the motor valves along with the main feed line on the Zipperclave. The feed was continued and there vessel pressure was regulated at 55 psi and a temperature of 1440°C. The automation was designed to hold the temperature and I the pressure within safe operating limits while

- 52 -

Page 54 0[ 11

addition of ethylene oxide proceeded through a pair of motor control valves. The feecd was allowed to continue until 60.0 grams of ethylene oxide (1.362 mobles) was added based on a difference weight of the feed cylinder. After the feed was complete, the reaction was allowed to continue for 1 hoour after which the vessel was cooled to 60°C, purged 4 times with nittrogen to 80 psi and was dumped to a container. The final product yield was 115 grams with a theoretical yield of 120 grams. The GPCC molecular weight of the product was Mw = 3550 and the MN = 22930 based on monodisperse polystyrene standards.

## Example 7

## Preparation of Poly(phenyl glycidyl ether)

To a 500 milliliter round bottom equipped with an overhead stirreer, nitrogen inlet, reflux condenser, addition funnel, and temperature controller was charged 47.06 grams of phenol (500 mmoles) and 1000 grams of toluene. The solution was brought to 50°C. Once heated, 1.00 milliliter (8 mmoles) of BF3/Et2O was added using a 2 milliliter syringe. Once the acid was added, 68.18 grams of phenyl glycidyl ether (4454 mmoles) was added dropwise so as to maintain a reaction temperature of 45°C-55°C. Once the glycidyl ether was added, the solution is refluxed for 3 hours, then cooled to about 50°C.

Whhile hot (<60°C) the organic was transferred to a separatory funnel and was washed once with 100 milliliters of 5% sodium bicarbonnate solution. The aqueous layer was drained and the organic was washed two more times with 100 milliliter portions of deionnized water. The aqueous layers were decanted and the organic was a dried for at least 1 hour over magnesium sulfate. Once dry the mnagnesium sulfate was filtered from the organic which was stripped of i solvent using a rotary evaporator. The final yield of viscous polymer was 90.3 grams (with 11% unreacted phenol). The GPC molecular 1 weight was Mw = 470 and the Mn = 310 (on average a trimer) based opn monodisperse polystyrene standards.

- 53 -

# Example 8 Preeparation of 1.3-Bis(phenoxy)-2-propanol uusing the Cascading Polvol Technique

To as 1 liter round bottom flask equipped with an overhead stirrer,r, nitrogen inlet, reflux condenser, addition funnel, and temperaturee controller was charged 94.11 grams of phenol (1 mole), 12.86 grams of tetraethylammonium iodide (0.05 moles), 3.00 grams of water (((0.17 moles), 42.08 grams of potassium hydroxide (0.75 moles), andd 250 grams of toluene. To a 100 milliliter additional funnel was charged 23.13 grams of epichlorohydrin (0.25 moles) and 50 grams of toluene. The solution was brought to 65°C at which time the epichlorohydrin solution was added over a period of 15 minutes while maintaining a reaction temperature of 65°C ± 5°C. The reaction was allowed to proceed for 48 hours.

Afteer 48 hours, the solution was cooled down to room temperature. The toluene solution was washed with two 250 milliliters portionns of deionized water. The aqueous layers were drained off, and the toluene was removed along with unreacted phenol using a rootary evaporator. The final yield of product was 64.5 grams which wass 106% of theory (residual is phenol). Final product purity was about t 95% as shown by GPC.

# Example 9 Dimnerization of 1.3-Bis(phenoxy)-2-propanol uusing the Cascading Polyol Technique

To an 250 milliliter round bottom flask equipped with an overhead stirrer,r, nitrogen inlet, reflux condenser, additional funnel, and temperature controller was charged 20.03 grams of 1,3-bis-

WO 93/24544

1 10 OL 206 1

(phenoxy)-2-proppanol prepared in Example 8 (82 mmoles), 2.06 grams of tetraethylammonium iodide (8 mmoles), 0.49 grams of water (27 mmoles), 6.51 grams of potassium hydroxide (116 mmoles), and 125 grams of toluenee. To a 100 milliliter addition funnel was charged 3.61 grams of eppichlorohydrin (39 mmoles) and 25 grams of toluene. The solution was brought to 65°C at which time the epichlorohydrin solution was addded over a period of 15 minutes while maintaining a reaction temperature of 65°C  $\pm$  5°C. The reaction was allowed to proceed for 48 hoours.

After 48 hours, the solution was cooled down to room temperature. The toluene solution was washed with two 250 milliliter portionns of deionized water. The aqueous layers were drained off, and I the toluene was removed using a rotary evaporator. The final yield obf product was 21.6 grams which was 101% of theory. GPC showed two major components of the product. The first was the starting material at about 41% (Mn = 220) and the second was the coupled product  $\epsilon$  at about 59% (Mn = 520).

# Example 10 Preparation of 1.3-Bis(hexadecyloxy)-2-propanol yusing the Cascading Polyol Technique

To a a 500 milliliter round bottom flask equipped with an overhead stirrerr, nitrogen inlet, reflux condenser, additional funnel, and temperature controller was charged 60.61 grams of hexadecanol (0.25 moles), 6.188 grams of tetraethylammonium iodide (0.024 moles), 1.44 grams of water (0.082 moles), 20.20 grams of potassium hydroxide (0.36 i moles), and 125 grams of toluene. To a 100 milliliter addition funnel i was charged 11.10 grams of epichlorohydrin (0.12 moles) and 25 grams of toluene. The solution was brought to 65°C at which time the epichlorohydrin solution was added over a period of

- 55 -

15 minutes while maintaining a reaction temperature of 65°C  $\pm$  5°C. The reaction was allowed to proceed for 48 hours.

Afteer 48 hours, the solution was cooled down to room temperature. The toluene solution was washed with two 250 milliliter portions of deionized water. The aqueous layers were drained off, and the toluene was removed using a rotary evaporator. The final yield off product was 70.9 grams which is 109% of theory (residual is hexaadecanol).

## Example 11

Sulfaation of 1.3-Bis(nonylphenoxy)-2-propanolblock-(http://propylene.oxide)10-block-(ethylene.oxide)10

To aa 250 milliliter round bottom flask equipped with an overhead stirrer,r, a temperature controller, and a vacuum adapter was added 75.0 ggrams of the material from Example 13 (49 mmoles). The kettle was thhen evacuated to <20 mmHg and heated to 100°C to remove any wateer. After 1 hour, the kettle was cooled to 60°C while under vacuum. 'When reaching 60°C, vacuum was broken with nitrogen and 5.33 grams of sulfamic acid (54 mmoles) was added. After charging thhe sulfamic acid, the kettle was heated to 110°C and evacuated to <200 mm Hg. The reaction was allowed to proceed for 3 hours.

At tithe end of the hold period, the kettle was cooled to 85°C and vacuum was broken with nitrogen. 1.2 grams of diethanolamine ((11 mmoles) was slowly added under a blanket of nitrogen. This scolution was stirred for 30 minutes. 10 grams of ethanol was added to the kettle and the temperature was regulated to 55°C. This solution was stirred for 30 minutes. The heat was removed from the kettle and 30 grams of water along with 20 grams of ethanol were  $\varepsilon$  added while maintaining good agitation. The

メガヘ コンドア・・・・・・・

- 56 -

solution was stirred for 15 minutes or until cooled to room temperature (<335°C).

Three pH was checked by dissolving 2 grams of the product solution in 18 ggrams of deionized water. If the pH was below 6.5, 0.2 gram increments of diethanolamine was added until the pH is between 6.5 and 0.5.

## Example 12

## FPreparation of 1.3-Bis(nonylphenoxy)-2propanol-block-(propylene oxide)<sub>10</sub>

To a 500 milliliter stainless steel Zipperclave was added 100.0 grams (0.2.202 moles) of 1,3-bis(nonylphenoxy)-2-propanol prepared in Example 1 along with 0.7 grams of potassium hydroxide. The vessel was attached to an automated unit and was heated to 50°C. The vessel was continuously purged with nitrogen for 15 minutes and waas then heated to 100°C where it was again continuously puurged with nitrogen for another 15 minutes. The vessel was then a heated to 140°C and is given a series of 6 purges by pressuring the v vessel up to 80 psi, and then venting. Once the venting process s was completed, the vessel was pressured to 20 psi with nitrogen.

Linnes connected to a cylinder which had been precharged withh 117.0 grams of propylene oxide (2.02 moles) were opened to the mnotor valves along with the main feed line on the Zipperclave. The feed was continued and the vessel pressure was regulated at 55 j psi and a temperature of 140°C. The automation was designed to hold the temperature and the pressure within safe operating limits: while addition of ethylene oxide proceeded through a pair of motor countrol valves. The feed was allowed to continue until all of the propylene oxide had been fed. After the feed was complete, the reaction was allowed to continue for 1 hour after which the vessel

Page 59 of 11

was cooled to  $60\,$ °C, purged 4 times with nitrogen to 80 psi and was dumped to a container. The final product yield was 211 grams with a theoretical yield 1 of 277 grams. The GPC molecular weight of the product was Mw v = 650 and the Mn = 490 based on monodisperse polystyrene stanndards.

## Example 13

## Preparation of f 1.3-Bis(nonylphenoxy)-2-propanol- block-(propylene oxide)<sub>10</sub>-block-(ethylene oxide)<sub>10</sub>

To aa 500 milliliter stainless steel Zipperclave was added 75.0 grams of thee propoxylate prepared in Example 12 (0.070 moles) along with 0.3 ggrams of potassium hydroxide. The vessel was attached to an auutomated ethoxylation unit and was heated to 50°C. The vessel was econtinuously purged with nitrogen for 15 minutes and was then heated to 100°C where it was again continuously purged with nitrogen for another 15 minutes. The vessel was then heated to 140°C \( \ext{c} \) and was given a series of 6 purges by pressuring the vessel up to 80 ppsi, and then venting. Once the venting process was completed, the veessel was pressured to 20 psi with nitrogen.

The e ethylene oxide lines were opened to the motor valves along with the mnain feed line on the Zipperclave. The feed was continued and thhe vessel pressure was regulated at 55 psi and a temperature of 1440°C. The automation was designed to hold the temperature andd the pressure within safe operating limits while addition of ethyldene oxide proceeded through a pair of motor control valves. The feedd was allowed to continue until 30.7 grams ethylene oxide (0.696 moldes) was added based on a difference weight of the feed cylinder. After tithe feed was complete, the reaction is allowed to continue for 1 hoour after which the vessel was cooled to 60°C, purged 4 times with nitrrogen to 80 psi and was dumped to a container. The

final product yield was 99 grams with a theoretical yield of 106 grams.

# Example 14 Preeparation of Bis(nonylphenoxy) Adduct of 1.4-Butanediol Diglycidyl Ether

To a five neck, two liter round bottom flask equipped with an addition funnnel, thermometer, nitrogen dispersant tube, mechanical stirrrer, and a decanting head with a water-cooled condenser were added 506.8 grams (2.30 mole) of nonylphenol and 350 milliliters of cyclohexane. The solution was heated to reflux, and 6.5 grams (1.3 weigght percent based on nonylphenol) of potassium hydroxide in 155 milliliters of water was slowly added to the round bottom flask. AAfter all the water was recovered in the decanting head (15 milliliters + + 2 milliliters formed), 220 grams (1.09 mole) of 1,4butanediol digly.ycidyl ether was added dropwise between 60 and 80°C. After the addition was complete, the solution was refluxed for four hours. The contitents of the flask were then washed with a five percent aqueouss solution of phosphoric acid, and the organic layer was separated fifrom the water layer and washed twice with deionized water. The reaction mixture was then placed in a one liter round bottom flask, annd the remaining cyclohexane and unreacted nonylphenol weere recovered by distillation, first at atmospheric pressure, then runder vacuum at 0.2 mm Hg. The kettle temperature was not allowedd to exceed 180°C during the distillation to prevent discoloration of f the product. The concentrated solution was then refiltered to give 710 grams of a pale-yellow liquid. Molecular weight by end-group MYW analysis was 689.9 (theoretical MW = 643.0). Ir and nmr spectra weere consistent with the expected structure of the product.

الإلاي بالماسات والما

- 59 -

# Example 15 Preparation of 3 Mole Ethoxylate of 1.3-Bis(nonylphenoxy)-2-propanol

To ) a five hundred milliliter Zipperclave reactor were charged, under r nitrogen, 200.1 grams (0.43 mole) of 1,3bis(nonylphenoxxy)-2-propanol prepared in Example 2 and 0.20 grams (0.1 weight percent) of BF3/Et2O. The reaction mixture was heated to 80°C, and 55.1 ggrams (1.25 mole) of ethylene oxide was fed to the reactor over a tywo hour period. After all the ethylene oxide was fed. the reaction mixxture was allowed to cook out for one hour and then dumped hot, unader nitrogen, into a jar containing 160 milliliters of a one percent aquueous solution of sodium hydroxide. The organic layer was separated fifrom the water layer and washed twice with deionized water. The washes were performed at 90°C to facilitate the separation of thee two layers. The product was then dried by azeotropic removval of the water, using cyclohexane (300 milliliters) as the entrainerr. The cyclohexane was stripped off under vacuum to give a pale-yellow liquid with a molecular weight by end-group MW analysis of 601.77 (theoretical MW = 629). Ir and nmr spectra were consistent with t the expected structure of the product.

# Example 16 Preparation of 8 Mole Ethoxylate of Bis(3(nonylphenoxy) Adduct of 1.4-Butanediol Diglycidyl Ether

To an five hundred milliliter Zipperclave reactor were charged, under nnitrogen, 150.2 grams (0.22 mole) of bis(nonylphenoxyy) adduct of 1,4-butanediol diglycidyl ether prepared in Example 14 annd 0.30 grams (0.2 weight percent) of BF3/Et2O. The reaction mixture 2 was heated to 80°C, and 77.5 grams (1.76 mole) of

- 60 -

ethylene oxide 1 was fed to the reactor over a two hour period. After all the ethylene oxxide was fed, the reaction mixture was allowed to cook out for one houur and then dumped hot, under nitrogen, into a jar containing 160) milliliters of a one percent aqueous solution of sodium hydroxide. Thee organic layer was separated from the water layer and washed twwice with deionized water. The washes were performed at 90°C to facililitate the separation of the two layers. The product was then dried by aazeotropic removal of the water, using cyclohexane (300 milliliters) as tithe entrainer. The cyclohexane was stripped off under vacuum to givee a pale-yellow liquid with a molecular weight by end-group MW analysis of 1047 (theoretical MW = 995). Ir and nmr spectra were consistent with the expected structure of the product.

### Examples 17 to 21

## Preparation of Propylene Oxide and Ethylene Oxide/Propylene Oxide CCopolymers of 1.3-Bis(nonylphenoxy)-2-propanol

To'o a 500 milliliter stainless steel autoclave was charged an amount of ppotassium hydroxide and starter listed in Table A below. The vessel was heated to 50° C. The vessel was continuously purged with nititrogen for 15 minutes, and was then heated to 100° C, where it was again continuously purged with nitrogen for another 15 minutes. The vessel was then heated to 140° C, and was given a series of 6 purgges by pressurizing the vessel up to 80 psi, and then venting. Once e the venting process was completed, the vessel was pressurized to 120 psi with nitrogen. The feed lines from an oxide feed cylinder (containing a charge of ethylene oxide or propylene oxide as identifed in Table A) to the autoclave were opened. The control system fed the e oxide at a rate such that the vessel pressure was regulated at 555 psi and a temperature of 140° C. The reaction was allowed to containue for 1 hour past the end of the oxide feed. The vessel was cooliled to 60° C, purged 4 times with nitrogen to 80 psi, and its contents we'ere dumped into a tared container. The final

WO 93/24544

PCT/US93/04872

- 61 -

theoretical and  $\epsilon$  actual product yields and molecular weights determined by ggel permeation chromatography are listed in Table A.

TABLEA

<del>e</del>					
смср	.005	.005	ပ	.04	.03
Surface Tension <sup>a</sup>	43.3	47.1	ပ	42.6	44.2
Mw	2340	3050	1530	2570	3060
Mn	2110	2720	1270	2280	2720
Actual yield	127	188	162	104	145
Theory yield	132	193	167	104	149
grams E0	82	145	0	54	66
grams PO	0	0	117	0	0
grams KOH	0.3	0.3	0.5	0.3	0.3
grams Starter	20	20	20	20	20
Starter Structure x y	10 0	0	0	0	0
Sta × Sta					
Product Structure x y	0 40	0 70	0	0 40	20 70
Example Product Structure x	17 1				

a - surface tension in dynes/cm at 1 wt.% copolymer in water

# b - approximate critical micelle concentration

# c - water insoluble

- 63 -

## Examples 22-33

## Preparation of Poly(nonvlphenyl glycidyl ether)

An 1 amount of ethanol listed in Table B below, and 200 grams of cyclohoexane were charged to a 500 milliliter round bottom flask equipped v with an overhead stirrer, nitrogen inlet, reflux condenser, additition funnel, and temperature controller. The solution was hexated to 50° C after which an amount of boron trifluoride etherrate listed in Table B was added. Subsequently, 100 grams of nonylpphenyl glycidyl ether was added dropwise to the reaction mixturge so as to maintain a reaction temperature of 45-55° C. The solutionn was refluxed at 83° C for an additional three hours after the completeion of feed, and then cooled to 50° C. While hot (<60° C), the organic 1 material was transferred to a separatory funnel, and was washed oncce with 100 milliliters of 5% sodium bicarbonate solution. The acqueous layer was washed two more times with 100 milliliter portionns of deionized water. The aqueous layers were decanted, and tithe organic layer was dried for at least 1 hour over magnesium sulflfate. Once dry, the magnesium sulfate was filtered from the organicic material, which was stripped of solvent using a rotary evaporator. The molecular weights of the polymer based on gel permeation chromatography are listed in Table B.

Table B

Example	mL <u>BF#3/Et2O</u>	Grams Ethanol	Mn	Мw
22	0.0.5	1.9	1700	· <b>260</b> 0
23	1.1.25	10.0	410	450
24	0.0.5	5.5	<b>47</b> 0	560
25	1.1.25	5.5	<b>87</b> 0	1150
<b>2</b> 6	1.1.25	1.0	<b>1580</b>	2530
27	22.0	5.5	900	1190
28	22.0	1.0	1470	2310
29	22.0	10.0	440	500

- 64 -

<u>Table B</u> (Continued)

Example	mL <u>BFf3/Et2O</u>	Grams Ethanol	Mn	Mw
30	00.5	10.0	580	730
31	00.5	1.0	1750	2790
32	00.5	1.0	<b>174</b> 0	2640
33	11.6	3.32	1170	<b>157</b> 0

## Examples 34-36

## Preparation of Unsymmetric Biphobes

To a 500 milliter round bottom flask equipped with an overhead stirreer, nitrogen inlet, reflux condenser, addition funnel, and temperature controller was charged an amount of starting alcohol listed inn Table C below. The material was heated to 170° C under nitrogen a sparge, after which an amount of potassium hydroxide listedd in Table C was added. An amount of epoxide listed in Table C was a fed to the reaction flask using an FMI feed pump at a rate such that addition of the epoxide was completed in 6 hours. Gel permeation chromatography showed that the purity of the biphobic product was greeater than 90%.

- 65 -

## Table C

$$egin{array}{c|c} R_{\overline{1}} & CH_2 \\ & CH & OH \\ R_{\overline{2}} & CH_2 \\ \end{array}$$

Ex.	Alcohol	GGms Al\lch.		Epoxide	Grams Epoxide	R1	<u>R2</u>
34	nonyl- phenol	10.00	2	1,2 epoxy- hexadecane	108.6	nonyl- phenoxy	tridecyl
35	dodecyl- phenol	10000	2	1,2 epoxy- hexadecane	91.1	dodecyl- phenoxy	tridecyl
<b>3</b> 6	dodecyl- phenol	50:0	1	nonylphenyl glycidyl ether	52.7	dodecyl- phenoxy	nonyl- phenoxy

## Example 37

## Prreparation of Poly(1.2-epoxyhexadecane)

Thee process described in Examples 34-36 was used with 50 grams of 1-haexadecanol, 1 gram of potassium hydroxide, and 49.4 grams of 1,2 epooxyhexadecane. The product multiphobe had a composition of 110% hexadecanol, 55.5% biphobe (i.e., 1-hexadecoxy-2-hexadecanol), 27.7.7% triphobe, and 6.5% quadphobe, as determined by gel permeation 1 chromatography.

## Example 38

Preparation c of an Unsaturated Poly(nonylphenyl glycidyl ether)

To a a 500 milliliter round bottom flask equipped with an overhead stirrerr, nitrogen inlet, reflux condenser, addition funnel, and temperature controller was charged 10 grams of allyl alcohol, 40

- 66 -

grams of toluenne, 0.5 grams of potassium hydroxide, and 2 grams of deionized water. The reaction mixture was refluxed to dryness, and cooled to 60° C.:. Once cool, 47.6 grams of nonylphenyl glycidyl ether was fed to the reaction vessel using a FMI pump over 35 minutes. The reaction minixture was heated to 112° C, and refluxed for 3 hours. The solvent was removed from the reaction mixture by vacuum stripping. The 3 product was cooled and recovered. The product composition wass 15% allyl alcohol, 43% biphobe (e.g., 1-(2-propene-1-oxy)-3-nonylphaenoxy-2-propanol) and 42% triphobe, as determined by gel permeationn chromatography.

## Example 39

## SSolventless Macromonomer Preparation

Too a 3 liter round bottom flask equipped with an overhead stirreer, nitrogen inlet and sparging tube, water cooled reflux condenseer, monomer addition tube, FMI pump and feed tank, and heating maantel and temperature controller, 2000 grams of previously meltited surfactant S-2 were charged. The materials were heated to 85° CC under nitrogen sparge and mixing, and held at temperature four 1 hour to drive off residual water. Then 0.05 grams of 4-methoxyphhenol were added, and the mixture was sparged with air for 15 minutites to activate the inhibitor. 2.4 grams of dibutyl tin dilaurate were added, and after 15 minutes of mixing, 201.25 grams of TMI were fedd over 45 minutes. The mixture was held at 85° C for another 4 hourss. Then 243 grams of water was pumped into the reaction mixture over a 25 minute period to wash the feed lines of isocvanate, andd to dilute the product macromonomer to 90% solids. The product maacromonomer was cooled and collected in a 1 gallon jug.

- 67 -

## Example 40

## Preparation of Macromonomer Compound

Intto a 1 liter round bottom reaction flask equipped with a heating mantle,; dean stark trap, condenser, thermometer, nitrogen bubbler, nitrogeen purge line and stirrer was charged 300 grams of toluene and 63 ggrams of a surfactant identified as S-1 in Table D below. With nititrogen purge, the resulting solution was heated to reflux at approxisimately 110°C and azeotroped to remove trace water to dryness. The esolution was subsequently cooled to 90°C, and 1.5 grams of bismutith hex chem 28% bismuth octoate catalyst (Mooney Chemical, Inc., (Cleveland, Ohio) was charged and allowed to mix well, after which a stoichiometric amount of 95% m-TMI aliphatic isocyanate (Ameerican Cyanamid, Stamford, Connecticut) was charged. After t the reaction proceeded at 90°C for 1.3 hours, the resulting produckt was cooled to 70°C and 0.03 grams of 2,6-di-tert-4methyl phenol (KBHT) preservative was added. The mixture was poured into a statainless steel pan with large surface area to facilitate drying. The finaal product was a waxy material, and is designated herein as macromonomer M-1.

## Table D

```
R_{1}\text{-O-CH}_{2}
|
CHH\text{-(OCH}_{2}CH_{2})_{x}OCH_{2}CH_{2}OH
|
R_{2}
R_{2} = \text{hyydrogen or a } R_{3}\text{-O-CH}_{2}\text{- residue}.
```

WO 93/24544

PCT/US93/04872

- 68 -

Surfactant	<u> </u>	R <sub>2</sub> /R <sub>3</sub>	Moles of Ethoxylation
S-1	No:onylphenol	Hydrogen (R <sub>2</sub> )	40
S-2	No:onylphenol	Nonylphenol (R <sub>3</sub> )	40
S-3	Noonylphenol	Nonylphenol (R3)	20
S-4	No:onylphenol	Octylphenol (R3)	20
S-5	No:onylphenol	Octylphenol (R3)	40
S-6	Noonylphenol	Nonylphenol (R <sub>3</sub> )	80
S-7	Noonylphenol	Nonylphenol (R3)	120
S-8	Noonylphenol	Nonylphenol (R3)	6
S-9	Noonylphenol	Nonylphenol (R3)	12

## Examples 41-61

## Proreparation of Macromonomer Compounds

In 1 a manner similar to that described in Example 40, other macromonomers were prepared using stoichiometric amounts of the surfactannts and unsaturated compounds identified in Table E below.

Table E

Example No.	Suurfactant	Unsaturated Compound	Macromonomer Designation
41	SS-2	m-TMI	M-2
42	SS-3	m-TMI	M-3
43	SS-4	m-TMI	M-4
44	SS-5	m- $TMI$	M-5
45	SS-6	$\mathbf{m}\text{-}\mathbf{TMI}$	<b>M-6</b>
46	SS-7	m- $TMI$	M-7
47	SS-2	Isocyanato Ethyl	M-8
		Methacrylate	

- 69 -

## <u>Table E</u> (Continued)

Example <u>No.</u>	Surrfactant		Macromonomer Designation
48	S-\$-5	Isocyanato Ethyl Methacrylate	M-9
49	S-;-1	Methacrylic Anhydrid	e <b>M-10</b>
50	S-3-2	Methacrylic Anhydric	de <b>M</b> -11
51	S-\$-5	Methacrylic Anhydrid	le M-12
52	S-6-6	Methacrylic Anhydric	de M-13
53	S-3-2	Acrylic Anhydride	M-14
54	S-5-5	Acrylic Anhydride	M-15
<b>5</b> 5	S-6-6	Acrylic Anhydride	M-16
56	S-2-2	Crotonic Anhydride	M-17
57	S-5-5	Maleic Anhydride	M-18
58	S-3-2	Methacryloyl Isocyan	ate M-19
59	S-6-6	Methacryloyl Isocyana	ate M-20
60	S-8-8	m-TMI	M-21
61	S-S-9	m-TMI	M-22

## Example 62

## Preparation of Alkali Soluble Thickener

A mmonomer mixture (300 grams) was prepared by charging ethyl aacrylate (Aldrich), methacrylic acid (Aldrich), macromonomer I M-1, 13 grams of a 75% solution of Aerosol OT surfactant (Ameerican Cyanamid, Stamford, Connecticut), and 3 grams of distilleed deionized water to a bottle, and dispersing the contents with vicigorous shaking. The ethyl acrylate, methacrylic acid and macromonomer M-1 were added in amounts identified in Table F below. A catalyst feed mixture comprised of 0.53 grams of sodium persulfate (Aldrich) and 52.47 grams of water was prepared in

- 70 -

another containner. To a 2 liter resin flask that had been immersed in a thermostated I water bath and equipped with a 4-bladed stainless steel mechanical stirrer, Claisen connecting tube, water condenser, nitrogen spargge and bubble trap, thermometer and monomer and catalyst addition inlets, 1.20 grams of the sodium salt of vinyl sulfonic acid and 658.5; grams of water were charged. The monomer mixture was charged to a 1-liter graduated monomer feed cylinder, and the catalyst solution was charged to a 125 milliliter graduated catalyst feed cylinder. Under nitrogen purge, the reactor was heated to 70°C. whereupon 33 r milliliters of the monomer mixture and 3 milliliters of the catalyst feeed mixture were charged to the reaction vessel. The reaction vessel 1 was subsequently heated to 80°C. After allowing the monomers to rejeact for 20 minutes to form a seed product, the monomer and catalyst feed mixtures were conveyed to the reaction vessel by FMI ppumps via 1/8" teflon tubing at a rate of 1.94 and 0.27 milliliters/n/minute, respectively, under continuous stirring at a reaction temperature held between 76-82°C. The reaction was allowed to proceed for another hour, after which the product was cooled and filterred with a 200 mesh nylon cloth. The coagulum was collected from tithe reaction vessel and filter cloth. Thickening ability of the resulting z product was monitored by Brookfield viscosity at 6 rpm by dilutingg the latex to 0.25%, 0.50% and 0.75% solids, and subsequently neeutralizing the product to pH=9.0 with a 95% solution of 2-amino-2-maethyl-1- propanol (AMP-95, Angus Chemical Company). Thee results are given in Table F.

## Examples 63-174

## PPreparation of Alkali Soluble Thickeners

In a manner similar to that described in Example 62, other alkali soluble thickeners were prepared using the monomers identified in Tables F-Q below in the amounts identified in Tables F-Q. Tables F and G illustrate the influence of m-TMI-containing

- 71 -

macromonomer (concentration and ethoxylation on thickening efficiency. Table e H illustrates the influence of 2-SEM on thickening efficiency. Tablde I illustrates the influence of mixing m-TMIcontaining macromonomers of various ethoxylations on thickening efficiency. Tablele J illustrates the influence of unsaturation type of urethane-containing macromonomers on thickening efficiency. Table K illustratites the influence of methacryloyl isocyanate incorporated into the macromonomer. Table L illustrates the influence of maccromonomer ester structure and ethoxylation on thickening efficiency. Table M illustrates the influence of acid type and concentration on thickening efficiency. Table N illustrates the influence of polyymer glass transition temperature and water solubility on thickening efficiency. Table O illustrates the influence of acrylamide inacorporated into the macromonomer. Table P illustrates the ininfluence of cross-linkable monomer concentration on thickening efficiency. Table Q illustrates the influence of mercaptan on thickening efffficiency. As used in Tables F-Q below, the following abbreviations haave the indicated meanings: MM = Macromonomer; EA = Ethyl Acryylate; MAA = Methacrylic Acid; AA = Acrylic Acid; MA = Methyl Accrylate; t-BA = t-Butyl Acrylate; n-BA = n-Butyl Acrylate; MMA == Methyl Methacrylate; 2-EHP = 2-Ethylhexyl Propionate Merccaptan; 2-SEM = 2-sulfoethyl methacrylate; and 2-HEA = 2-Hydroxxy Ethyl Acrylate.

		Thickener C	Thickener Composition by Weight	n by	Brookfie	Brookfield Viscosity (CPS) @ nH=9 0		Ę.
Example	Macromonomer	%WW	%EA	%MAA	0.25%	0.50%	0.75%	Designation
G	M-1	10	20	40	06	380	1.000	F.4
2 E	1 676 Z	ren	וכוני וכוני	40	270 270	11 400	103 600	, <u>0</u> ,0
64	7-7- W-2	9 01	20	40	120	3,100	60.000	7-7 7-7-7-1
* 14 5 4	M-2	10	20	40	105	10,400	130,000	P-3a
	M-2	20	40	40	25	2,150	50,500	P-4
67	M-2	30	30	40	10	790	20,000	P-5
. œ	M-3	D	55	40	390	2,260	17,900	P-6
8 6	W-3	6.5	53.5	40	142	1,200	18,500	P-7
2 6	M	10	50	40	220	3,050	40,000	P-8
7.1	M-3	20	40	40	75	2,350	27,500	P-9
5.	M.4	10	20	40	242	4,400	39,000	P-10
73	M-5	10	20	40	45	7,400	84,000	P-11
2.5	N.	20	40	40	34	4,450	29,000	P-12
- F	M-6	rc	55	40	460	25,500	88,000	P-13
9.5	9-W	10	20	40	105	39,000	150,000	P-14
2.5	9-W	15	45	40	195	43,000	140,000	P-15
- 02	M.6	20	40	40	125	52,500	187,000	P-16
0 6	9-W	30	30	40	315	26,500	162,000	P-17
£ 0	M.7	rc	55	40	230	7,800	15,800	P-18
00 10	M-7	10	20	40	006	17,400	35,000	P-19
1								

TABLEG

-73-

		Thicken	Thickener Composition by	on by	F			
Example	Macromonomer	%WM	weignt %EA	%MAA	Dro 0.25%	Drookfield Viscosity (CPS) @ pH=9.0 0.5% .75%	(CPS) @ pH=9.0 .75%	1%
83	M-21 M-22	10 10	50	40	120 122	117 190	110 335	125 950
				H	TABLEH			
Example	Macromonomer	Thicken	Thickener Composition by Weight %MM %EA %	on by %MAA	Bro % 2-SEM	Brookfield Viscosity (CPS) @ pH=9.0 M 0.25% 0.5%	(CPS) @ pH=9.0 0.5%	.75%
84 .	M-6 M-6	10 10	50 50	35 30	5 10	23 <10	4,340 510	32,000 6,400

-74-

					TABLE				
Example	Macromonomer	Thickene %MM	Thickener Composition by Weight %MM %EA %	nn by %MAA	0.25%	Brookfield Viscosity (CPS) @ pH=9.0 0.5%	(CPS) @ pH=9.0 .75%	Thickener Designation	
986	M-3:M-6 1:1	10	50	40	225	24,000	85,000	P-20	1
87	M-2:M-6 1:1	10	50	40	135	21,200	134,000	P-21	
					TABLEJ				
		Thicken	Thickener Composition by Weight	ın by		Brookfield Viscosity (CPS) @ nH-9 0	(CPS) @ nH-9 0		
Example	Macromonomer	%WW	%EA	%MAA	0.25%	0.5%	.75%	Designation	
88	8 <del>-</del> 2	ល	52	40	250	14,800	124,000	P-22	
	M-8	10	20	40	93	11,200	125,400	P-23	
06	M-8	20	40	40	45	6,140	84,500	P-24	
91	M.9	2	55	40	275	6,200	27,000	P-25	
26	6-W	10	20	40	220	10,100	80,000	P-26	
93	M-9	20	40	40	<u>6</u>	7,800	000'06	P-27	
76	M-9	30	30	40	45	5,200	000'69	P-28	

$\mathbf{x}$
Œ
ABLE
4
⊻
L

. •		Thickene	Thickener Composition	<u></u>				
Example	Macromonomer	%MM	Weight %EA	"%MAA	Brookfield 0.25%	Brookfield Viscosity (CPS) @ pH=9.0 0.25% 0.5%	3) @ pH=9.0	
95	M-19	ເລ	55	40	167	4,050	28.000	
96	M-19	10	20	40	652	22,000	88,500	
	M-19	20	40	40	1,710	34,200	128,000	
. 00	M 10	76	Ç	4	0,404	004,40	00000	
86	M-19	30	30	40	2,125	61,300	155,000	
66	M-20	2	52	40	344	10,960	30,300	
100	M-20	10	20	40	1,115	29,000	68,500	
101	M-20	20	40	40	2,175	45,500	129,000	
102	M-20	30	30	40	2,400	61,000	150,000	

-76-

	-			Table L	Ţ			
	-	Thickene	Thickener Composition by	n by		Brookfield Viscosity	ęţ.	
Example	Macromonomer	%WW	%EA	%MAA	0.25%	(CF3) & pn=3.0 0.50%	0.75%	Thickener Designation
103	M-10	10	20	40	130	285	410	0.90
104	M.11	10	20	40	190	19.500	152 000	67. T
105	M-11	8 8	40	40	120	13,500	146,000	F-30
106	M-11	30	30	40	96	8,000	73.000	P-32
107	M.19	IC IC	ፎ ፎ ር	çç	980	2 5 5 7 Y	11.000	2 6 6
707	21-171 21-17	ء ج	8 5	2 5	175	0,4,0	01,000	F-33
108	M-12	2 6	8 <del>4</del>	40	0.01	7,400	77,000	P-34
110	M.19	e S	e S	40	62	4.500	000,11	r-00 D 26
111	M-13	വ	32	40	320	25,600	79,000	F-30
112	M-13	01	20	40	97	28,000	125,000	P.38
113	M-13	8	40	40	300	58,200	171,000	P-39
114	M-13	30	30	40	730	63,000	163,000	P-40
115	M-14	10	20	40	410	22,700	130,000	P-41
116	M-14	20	40	40	1225	44,500	168,000	P-42
117	M-14	30	30	40	1010	42,500	180,000	P-43
118	M-15	20	55	40	84	1,680	29,000	P-44
119	M-15	10	20	40	350	12,000	83,000	P-45
120	M-15	20	40	40	220	24,500	122,000	P-46
121	M-15	30	30	40	1050	33,000	133,000	P-47
122	M-16	വ	55	40	420	17,720	45,300	P-48
123	M-16	10	20	40	1,345	27,000	98,000	P-49
124	M-16	20	40	40	3,450	65,800	158,000	P-50
125	M-16	30	30	40	11,600	81,000	157,000	P-51
126	M-17	10	20	40	410	12,000	000'09	P-52
127	M-17	20	40	40	255	10,600	46,300	P-53
128	M-17	30	30	40 ·	38	2,525	13,500	P-54
129	M-18	ល	55	40	100	810	3,500	P-55
130	M-18	10	20	40	110	1,420	5,940	P-56
131	M-18	20	40	40	30	870	2,425	P-57

-77-

	Thickener % Designation					10-1 10-1 10-1 10-1 10-1 10-1 10-1 10-1								
	0.75%	1001	50.000	3	œ	43 500		8	79.0	28.0	52.5	122.	73	66 000
	Brookfield Viscosity (CPS) @ pH=9.0 .25% 0.50%	12.200	3.800	<10	95	13.500	13,000	24,000	7,600	7,060	16,700	28,000	213	16,500
	Brookf (CPS) 0.25%	1520	45	<10	<10	ਦਿ	210	7.7	17	130	86	130	<10	710
M	%AA	0	0	0	30	00	0	0	0	0	0	0	20	20
Table M	Thickener Composition by Weight MM %EA %MAA	30	20	10		88	30	30	30	20	20	20	0	20
	ner Compos %EA	99	70	80	09	<u></u>	65	09	20	45	40	30	70	20
•	Thicke %MM	10	10	10	10	200	ın.	10	20	2	10	20	10	10
	Macromonomer	M-2	M-2	M-2	M-2	M-6	M-6	9-W	9-W.	9-W	M-6	9-W	M-11	M-17
	Example	132	133	134	135	136	137	138	139	140	141	142	143	144

-78-

Example	Macromonomer	Thickener ( %MM	Thickener Composition by Weight %MM %EA %MAA	by Weight %MAA	%Other	0.25%	Brookfield Viscosity (CPS) @ pH=9.0 0.50%	0.75%	Thickener Designation
145	M-2	10	40	40	10 MMA	06	5,760	82,000	P-71
146	M-2	10	30	40	20 MMA	15	1,125	55,000	P-72
147	M-2	10	50 70 70	40	30 50 MMA	22	207	0,000	P-73
148	M-2	10	0	40	50 MMA	<10	<10	<10	P-74
149	M-2	10	30	40	20 styrene	20	310	1,330	P-75
150	M-2	10	40	40	10 styrene	92	7,540	75,500	P-76
151	M-2	10	40	40	10 n-BA	220	13,800	118,000	P-77
152	M-2	10	30	40	20 n-BA	185	7,400	66,500	P-78
153	M-2	10	40	40	10 t-BA	130	10,100	100,000	P-79
154	M-2	10	30	40	20 t-BA	125	7,200	77,500	P-80
155	M-2	10	40	40	10 MA	100	006'9	121,000	P-81
156	M-2	10	30	40	20 MA	73	5,000	90,000	P-82
157	M-6	20	30	40	10 MMA	83	15,400	150,000	P-83

	pH=9.0 .75%	162 124,000 110,000 104,000
	Broofield Viscosity (CPS) @ pH=9.0 0.25% 0.5%	137 32,700 22,500 22,300
	Broofield 0.25%	102 112 71 115
le Q	%Acryl- amide*	10 10 5 0
Table O	ion by %MAA	35 55 55 55 55 55 56 55 55 55 55 55 55 55 55 55 55 55 55 5
	Thickener Composition by Weight %MM %EA %	55 45 50 55
	Thickene %MM	0 10 10 10
	Macromonomer	none M-6 M-6
-	Example	158 159 160 161

\*The amount of acrylamide influences the pH of solubilization for the polymer. The pH of solubility for the polymer is as follows: Example 159 = 5.0; Example 160 = 5.8; and Example 161 = 6.2. The pH of solubility is that pH where the aqueous sample goes from being a low viscosity and turbid latex to a viscous clear solution.

-79-

Table P

# PCT/US93/04872

WO 93/24544
-------------

-80-
------

Example	Macromonomer	I nickener %MM	Composition WEA	1 nickener Composition by weight %MM %EA %MAA	%2-HEA	0.25%	(CPS) @ pH=9.0 0.50%	0.75%	Thickener Designation
162	M-2	10	47.7	40	2.3	97	090'6	127,000	P-84
163	7-M	Q (2	 	8 €	9 6 U	20 6	6,300	76,000	P-85
104	7-W	8 8		Q# <b>9</b>		77	6,200	116,600	P-86
165	Z-W	0 20	5 5 7	9.0	o t	01°	260	18,600	P-87
991	M-2	S 8	32.5	940	d.7.	20	720	40,000	P-88
167	M-2	20	30	40	10	10	520	29,500	P-89
				II	Table Q				
•		Thickener	. Compositic	Thickener Composition by Weight	*dng 6 <i>%</i>	i d	Brookfield Viscosity (CPS) @ pH=9.0		Thickener
Example	Macromonomer	WIMIW.	70EA	%INIAA	702-Enr	0.25%	0.50%	0.75%	Designation
168	M-2	10	40	20	.05	165	22,800	142,000	P-90
169	M-2	10	20	40	0.5	18	2,060	66,500	P-91
170	M-2	10	20	40	0.3	<10	115	9,700	P-92
171	M-2	10	20	40	0.5	<b>~10</b>	12	355	P-93
172	M-2	10	20	40	<b>-</b>	<b>&lt;1</b> 0	<10	<10	P-94
173	9-W	10	20	40	.05	230	23,700	90.700	P.95
174	9-W	10	20	40	63	90	5.170	33 000	90 €

\*% charged to reactor based on monomer.

WO 93/24544

PCT/US93/04872

- 81 -

#### **Examples 175-230**

## Co-Thickening wwith Surfactants

The e addition of certain surfactants to an associative polymer solution produces a co-thickening effect. The results in Table L below shhow the co-thickening effect produced by the addition with thorough mnixing of certain surfactants identified in Table R below in the amounts identified in Table L to a 0.5% alkaline solution of an alkali solubble thickener identified in Table S as measured with a Brookfield Viscometer at 6 rpm at pH = 9.0.

### Table R

$$R_1$$
-O-CH<sub>2</sub>

| CH-(OCH<sub>2</sub>CH<sub>2</sub>)<sub>x</sub>OCH<sub>2</sub>CH<sub>2</sub>OH

| R<sub>2</sub>
 $R_2$  = hydrogen or a  $R_3$ -O-CH<sub>2</sub>- residue.

Surfactant	R <sub>1-1</sub>	<u>R<sub>2</sub>/R<sub>3</sub></u>	Moles of Ethoxylation
S-8	Nonnylphenol	Nonylphenol (R <sub>3</sub> )	20
S-9	Nonnylphenol	Nonylphenol (R <sub>3</sub> )	40
S-10	Nonnylphenol	Nonylphenol (R <sub>3</sub> )	80
S-11	Nonnylphenol	Hydrogen (R <sub>2</sub> )	25
S-12	Nonnylphenol	Hydrogen (R <sub>2</sub> )	40
S-13	Nonnylphenol	Octylphenol (R3)	20
S-14	Nonnylphenol	Octylphenol (R <sub>3</sub> )	<b>4</b> 0
S-15*	Nonnylphenol	Nonylphenol (R3)	40
S-16	Octylyphenol	Hydrogen (R <sub>2</sub> )	25

- 82 -

# \* Sulfated derivative.

Table S

		Surfactan		n 16 11 Tr
		Concentra		Brookfield Viscosity
Example	Surfactarant	(wt.%)	Thickener	(cps) @ pH=9.0
175	S-8 3	0.0	P-3	3100
2.0	S-8 3	0.2	P-3	32700
	S-8 3	0.4	P-3	45700
	S-8.3.	0.8	P-3	63300
	S-8 3	1.0	P-3	65500
	S-8 3	2.0	P-3	>100000
	5-0 7	2.0		2100000
176	S-9 )	0.2	P-3	24200
	S-9 )	0.4	P-3	18700
	S-9 )	0.8	P-3	6600
	S-9 )	1.0	P-3	4060
		2.0	P-3	<b>122</b> 5
177	S-100	0.2	P-3	20600
	S-100	0.4	P-3	17300
	S-100	0.8	P-3	8500
	S-100	1.0	P-3	<b>6300</b>
	S-100	2.0	P-3	1850
			5.0	1000
178	S-1111	0.2	P-3	12000
	S-1111	0.4	P-3	3160
	S-1111	0.8	P-3	700
	S-1111	1.0	P-3	485
	S-1111	2.0	P-3	480
179	S-1212	0.2	P-3	9200
2.0	S-1212	0.4	P-3	4500
	S-1212	0.8	P-3	1000
	S-1212	1.0	P-3	. 875
	S-122	2.0	P-3	565
	<del></del>			
180	S-133	0.2	: P-3	34300
	S-133	0.4	P-3	26700
	S-133	0.8	P-3	11500
• .	S-133	1.0	P-3	8600
	S-1313	2.0	P-3	2450

WO 93/24544

PCT/US93/04872

- 83 -

<u>Table S</u> (Continued)

		Surfactani Concentra		Brookfield Viscosity
Example	Surfactarant	(wt.%)	Thickener	(cps) @ pH=9.0
181	S-14 4	0.2	P-3	22200
	S-14 4	0.4	P-3	17200
	S-14 4	0.8	P-3	6900
	S-14 4	1.0	P-3	4500
	S-14 4	2.0	P-3	1500
182	S-15 5	0.2	P-3	10500
	S-15 5	0.4	P-3	4940
	S-15 5	0.8	P-3	2160
	S-15 5	1.0	P-3	1450
	S-15 5	2.0	P-3	355
183	S-16 3	0.2	P-3	14300
	S-16 3	0.4	P-3	4080
	S-16 3	0.8	P-3	1075
	S-16 3	1.0	P-3	735
	S-16 3	2.0	P-3	4.85
184	S-8	0.0	P-2	11400
	S-8	0.2	P-2	23500
	S-8	0.4	P-2	34000
	S-8	0.8	P-2	64000
	S-8	1.0	P-2	71000
	S-8	2.0	P-2	93000
185	S-9	0.2	P-2	11000
	S-9	0.4	P-2	4000
	S-9	0.8	P-2	2000
	S-9	1.0	P-2	1400
	S-9	2.0	P-2	850
186	S-10 )	0.2	P-2	10500
	S-10 )	0.4	P-2	5000
	S-10 )	0.8	P-2	2000
	S-10 )	1.0	P-2	1600
	S-10 )	2.0	P-2	950

- 84 -

Table S (Continued)

		Surfactant Concentrat		Brookfield Viscosity
T	Surfactcant	(wt%)	Thickener	(cps) @ pH=9.0
Example	Sunacidant	(WL-70)	Tinckener	(CDS) & DIX-V.V
187	S-3-11	0.2	P-2	2700
	S-3-11	0.4	P-2	1000
	S-3-11	0.8	P-2	800
	S-3-11	1.0	P-2	660
	S-3-11	2.0	P-2	620
188	S-3-12	0.2	P-2	2800
200	S-3-12	0.4	P-2	1000
	S-3-12	0.8	P-2	850
	S-3-12	1.0	P-2	660
	S-3-12	2.0	P-2	650
189	S-3-8	0.0	P-4	2150
	S-3-8	0.2	P-4	19000
	S-3-8	0.4	P-4	31000
	S-3-8	0.8	P-4	55000
	S-3-8	1.0	P-4	61000
	S-3-8	2.0	P-4	85000
190	S-3-9	0.2	P-4	19500
	S-3-9	0.4	P-4	21500
	S-3-9	0.8	P-4	11500
	S-3-9	1.0	P-4	7400
	S-3-9	2.0	P-4	2250
191	S-S-10	0.2	P-4	12600
	S-S-10	0.4	P-4	17400
	S-S-10	0.8	P-4	12600
	S-S-10	1.0	P-4	6600
	S-S-10	2.0	P-4	2600
192	S-S-11	0.2	P-4	17400
	S-S-11	0.4	P-4	7800
	S-S-11	0.8	P-4	1650
	S-S-11	1.0	P-4	860
	S-S-11	2.0	P-4	560

WO 93/24544

PCT/US93/04872

- 85 -

<u>Table S</u> (Continued)

		Surfactant	t	
		Concentra	tion	Brookfield Viscosity
Example	Surfactanınt	(wt.%)	Thickener	(cps) @ pH=9.0
193	S-12 2	0.2	P-4	14600
	S-12 2	0.4	P-4	7800
	S-12 2	0.8	P-4	1500
	S-12 2	1.0	P-4	960
	S-12 ?	2.0	P-4	450
194	S-8	0.0	P-5	790
	S-8	0.2	<b>P</b> -5	4600
	S-8	0.4	P-5	19600
	S-8	0.8	<b>P-</b> 5	42000
	S-8	1.0	P-5	50000
	S-8	2.0	P-5	90000
195	S-9	0.2	P-5	5800
	S-9	0.4	P-5	13200
	S-9	0.8	P-5	9200
	S-9	1.0	<b>P-5</b>	5200
	S-9	2.0	P-5	1600
196	S-10 )	0.2	<b>P-</b> 5	4050
	S-10)	0.4	<b>P-</b> 5	10400
	S-10)	0.8	<b>P-</b> 5	9400
	S-10 )	1.0	P-5	5000
	S-10 )	2.0	P-5	1600
197	S-11 l	0.2	<b>P-</b> 5	10600
	S-11 l	0.4	P-5	4200
	S-11 l	0.8	<b>P</b> -5	1400
	S-11 t	1.0	P-5	970
	S-11 i	2.0	<b>P-</b> 5	410
198	S-12 :	0.2	<b>P-</b> 5	6000
	S-12 ?	0.4	P-5	4200
	S-12 }	0.8	P-5	1150
	S-12 ?	1.0	P-5	600
	S-12 ?	2.0	P-5	· 340

WO 93/24544

PCT/US93/04872

- 86 -

Table S (Continued)

		Surfactan	t	•
		Concentra		Brookfield Viscosity
Example	Surfactavant	(wt,%)	Thickener	(cps) @ pH=9.0
199	S-8 3	0 '	P-7	1200
	S-8 3	0.2	P-7	9000
	S-8 3	0.4	P-7	21000
	S-8 }	0.8	P-7	37000
	S-8 3	1.0	P-7	49000
	S-8 3	2.0	P-7	78000
200	S-9 )	0.2	P-7	1600
	S-9 }	0.4	P-7	1350
	S-9 )	0.8	P-7	900
	S-9 )	1.0	P-7	762
	S-9 )	2.0	P-7	565
201	S-100	0.2	P-7	1100
	S-100	0.4	P-7	1150
	S-100	0.8	P-7	900
	S-100	1.0	P-7	823
	S-100	2.0	P-7	650
202	S-1111	0.2	P-7	1175
	S-1111	0.4	P-7	<b>68</b> 5
	S-1111	0.8	P-7	503
	S-1111	1.0	P-7	495
	S-1111	2.0	P-7	502
203	S-122	0.2	P-7	950
	S-122	0.4	P-7	<b>67</b> 5
	S-1212	0.8	P-7	<b>52</b> 5
	S-122	1.0	P-7	500
	S-1212	2.0	P-7	480
204	S-8 3	0.0	P-13	25500
	S-8 3	0.2	P-13	31500
	S-8 3	0.4	P-13	46500
	S-8 3	0.8	P-13	60000
	S-8 3	1.0	P-13	60000
	S-83	2.0	P-13	62500
	·			

- 87 -

Table S (Continued)

		Surfactan		Prochecia	Winnester
	0.6.4.4	Concentra		Brookfield	
Example	Surfactantnt	(wt.%)	<u>Thickener</u>	(cps) @ pH:	=9.0
205	S-9	0.2	P-13	8640	
	S-9	0.4	P-13	2940	
	S-9	0.8	P-13	1200	
	S-9	1.0	P-13	1000	
	S-9	2.0	P-13	<b>75</b> 0	
206	S-10 )	0.2	P-13	10100	
	S-10 )	0.4	P-13	4200	
	S-10 )	0.8	P-13	1450	
•	S-10)	1.0	P-13	1300	
	S-10 )	2.0	P-13	900	
207	S-12 !	0.2	P-13	2540	
	S-12 !	0.4	P-13	1125	
	S-12 :	0.8	P-13	750	
	S-12 !	1.0	P-13	670	
	S-12 !	2.0	P-13	610	
208	S-8	0.0	P-14	39000	
	S-8	0.2	P-14	61000	
	S-8	0.4	P-14	73500	
	S-8	0.8	P-14	87000	
	S-8	1.0	P-14	93500	
	S-8	2.0	P-14	122000	
209	S-9	0.2	P-14	41000	
	S-9	0.4	P-14	13700	
	S-9	0.8	P-14	6200	
	S-9	1.0	P-14	3500	
	S-9	2.0	P-14	1200	
210	S-10 )	0.2	. P-14	38200	
	S-10 )	0.4	P-14	20500	
	S-10 )	0.8	P-14	7300	
	S-10)	1.0	P-14	5400	
	S-10 )	2.0	P-14	1950	

- 88 -

Table S (Continued)

	D 6 4	Surfactant Concentrat	tion	Brookfield (cps) @ pH	
Example	Surfactanint	(wt.%)	Thickener	(cds) w bri	=3.0
211	S-12 2	0.2 ·	P-14	13000	
	S-12 2	0.4	P-14	4300	
	S-12 2	0.8	P-14	975	
	S-12 2	1.0	P-14	950	
	S-12 2	2.0	P-14	660	
212	S-8	0.0	P-16	52500	
	S-8	0.2	P-16	95000	
	S-8	0.4	P-16	92000	
	S-8	0.8	P-16	122000	
	S-8	1.0	P-16	125000	
	S-8	2.0	P-16	138000	
213	PS-9.9	0.2	P-16	73500	
	PS-9-9	0.4	P-16	53000	
	PS-9.9	0.8	P-16	25000	
	PS-99	1.0	P-16	21000	
	PS-9-9	2.0	P-16	5400	
214	S-10 0	0.2	P-16	52800	
	S-10 0	0.4	P-16	34500	
	S-10 0	0.8	P-16	5400	
	S-10 0	1.0	P-16	<b>292</b> 5	
	S-10 0	2.0	P-16	<b>7</b> 75	
215	S-13 3	0.2	P-16	45800	
	S-13 3	0.4	P-16	54000	
	S-13 3	0.8	P-16	<b>50800</b>	
	S-133	1.0	P-16	<b>5450</b> 0	
	S-13 3	2.0	P-16	63000	
216	S-144	0.2	P-16	22700	
	S-14 4	0.4	P-16	2480	
	S-14 4	0.8	P-16	710	
	S-144	1.0	P-16	532	
	S-14 4	2.0	P-16	415	

- 89 -

<u>Table S</u> (Continued)

		Surfactant Concentra		Brookfield Viscosity
Example	Surfactantnt	(wt.%)	Thickener	(cps) @ pH=9.0
Datimple	DOMESTIC	1114/2/		1000/ 0 044-0.0
217	S-8	0.0	P-29	285
	S-8	0.2	P-29	285
	S-8	0.4	P-29	360
	S-8	0.8	P-29	477
	S-8	1.0	P-29	505
	S-8	2.0	P-29	837
218	S-9	0.2	P-29	282
	S-9	0.4	P-29	285
	S-9	0.8	P-29	284
	S-9	1.0	P-29	298
	S-9	2.0	P-29	322
219	S-10	0.2	P-29	272
	S-10	0.4	P-29	278
	S-10	0.8	P-29	285
	S-10	1.0	P-29	<b>297</b>
	S-10	2.0	P-29	315
220	S-12	0.2	P-29	267
	S-12	0.4	P-29	279
	S-12	0.8	P-29	298
	S-12	1.0	P-29	311
	S-12	2.0	P-29	320
221	S-8	0.0	P-30	19500
	<b>S-</b> 8	0.2	<b>P-3</b> 0	<b>7900</b> 0
	S-8	0.4	P-30	71200
	S-8	0.8	P-30	81000
	S-8	1.0	P-30	89500
	S-8	2.0	P-30	175000
222	S-9	0.2	<b>P</b> -30	52000
	S-9	0.4	P-30	35500
	S-9	0.8	P-30	16500
	S-9	1.0	P-30	15600
	S-9	2.0	P-30	5620

WO 93/24544

PCT/US93/04872

- 90 -

Table S (Continued)

		Surfactant Concentra	tion	Brookfield Viscosity
Example	Surfactatant	(wt.%)	Thickener	(cps) @ pH=9.0
223	S-100	0.2	P-30	47200
	S-100	0.4	P-30	26300
	S-100	0.8	P-30	20300
	S-1Q0	1.0	P-30	13400
	S-100	2.0	P-30	4700
224	S-1712	0.2	P-30	23000
	S-1712	. 0.4	P-30	6840
	S-1712	0.8	P-30	<b>312</b> 5
	S-1712	1.0	P-30	1750
	S-1712	2.0	P-30	1225
225	S-88	0.0	P-46	24500
	S-88	0.2	P-46	79000
	S-88	0.4	P-46	75000
	S-88	0.8	P-46	86000
	S-88	1.0	P-46	95000
	S-88	2.0	P-46	150000
226	. S-99	0.2	P-46	40500
	S-9 9	0.4	P-46	31000
	S-99	0.8	P-46	15300
	S-9 9	1.0	P-46	9400
	S-9 9	2.0	P-46	2300
227	S-1:11	0.2	P-46	20000
	S-1:11	0.4	P-46	7300
	S-1:11	0.8	<b>P-4</b> 6	1350
	S-1:11	1.0	P-46	900
	S-1:11	2.0	P-46	380
228	S-1:13	0.2	P-46	63500
	S-1:13	0.4	P-46	42000
	S-1:13	0.8	P-46	23000
	S-1:13	1.0	P-46	16000
	S-1;13	2.0	P-46	4850

- 91 -

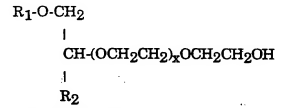
<u>Table S</u> (Continued)

		Surfactant Concentration		Brookfield Viscosity
Example	Surfactanint	(wt.%)	Thickener	(cps)@pH=9.0 `
229	S-14 !	0.2	P-46	36000
	S-14 I	0.4	P-46	25000
	S-14	0.8	P-46	11000
	S-14 l	1.0	P-46	9300
	S-14 !	2.0	P-46	1900
230	S-16 }	0.2	P-46	19000
	S-16 ;	0.4	P-46	9300
	S-16 }	0.8	P-46	<b>125</b> 0
	S-16 )	1.0	P-46	750
	S-16 }	2.0	P-46	290

#### **Examples 231-275**

#### Co-Thickening wwith Surfactants

The edgree of ethoxylation of a surfactant added to an associative polymer solution influences the co-thickening effect. The results in Table I U below show the co-thickening effect produced by the addition withh thorough mixing of certain surfactants identified in Table T below in 1 the amounts identified in Table U to a 0.3% (Examples 172-1889), 0.5% (Examples 190-215) or 0.75% (Example 216) alkaline solution 1 of an alkali soluble thickener identified in Table U as measured with a Brookfield Viscometer at 6 rpm at pH = 9.0.



 $R_2$  = hydrogen or a  $R_3$ -O-CH<sub>2</sub>- residue.

# Table T

Surfactant	RR_1	R2/R3	Moles of Ethoxylation
S-17	Nonnylphenol	Hydrogen (R <sub>2</sub> )	4
S-18	Nonnylphenol	Hydrogen (R <sub>2</sub> )	6
S-19	Nonnylphenol	Hydrogen $(R_2)$	7
S-20	Nonnylphenol	Hydrogen (R <sub>2</sub> )	8
S-21	Nonnylphenol	Hydrogen $(R_2)$	9
S-22	Nonnylphenol	Hydrogen (R <sub>2</sub> )	10
S-23	Nonnylphenol	Hydrogen $(R_2)$	15
S-24	Nonnylphenol	Hydrogen (R <sub>2</sub> )	25
S-25	Nopnylphenol	Hydrogen (R2)	<b>4</b> 0
S-26	Octatylphenol	Hydrogen (R <sub>2</sub> )	1
S-27	Octatylphenol	Hydrogen (R <sub>2</sub> )	3
S-28	Octatylphenol	Hydrogen (R <sub>2</sub> )	5
S-29	Octylphenol	Hydrogen (R <sub>2</sub> )	7
S-30	Octitylphenol	Hydrogen (R <sub>2</sub> )	9
S-31	Octitylphenol	Hydrogen (R <sub>2</sub> )	12
S-32	Octitylphenol	Hydrogen $(R_2)$	16
S-33	C111-C15	Hydrogen (R <sub>2</sub> )	5
S-34	Seccondary Alccohol C111-C15 Seccondary Aldcohol	Hydrogen (R <sub>2</sub> )	9

Table U

		Surfactar Concentr		Brookfield Viscosity
Example	Surfaactant	(wt.%)		$\frac{\text{(cps) @ pH = 9.0}}{\text{(cps) @ pH = 9.0}}$
Example	Dutteament		THICKELLET	_(cps/ & pii = 3.0
231	S-177	0.8	P-1	890
232	S-188	0.8	P-1	1340
233	S-199	0.8	P-1	<b>63</b> 0
234	S-200	0.8	P-1	205
235	S-211	0.8	P-1	143
236	S-222	0.8	P-1	113
237	S-233	0.8	P-1	85
238	S-24 4	0.8	P-1	57
239	S-25	0.8	P-1	68
240	S-17	0.8	P-3	17800
241	S-18 ::	0.8	P-3	35800
242	S-19	0.8	P-3	21300
243	<b>S-2</b> C	0.8	P-3	820
244	S-2	0.8	P-3	. <b>23</b> 0
245	S-2	0.8	P-3	147
246	S-23	0.8	P-3	118
247	S-24	0.8	P-3	82
<b>24</b> 8	<b>S-2</b> 5 ≎	8.0	P-3	<b>7</b> 7
249	S-177	8.0	P-42	<b>5700</b> 0
250	<b>S-18</b> 8	8.0	P-42	134000
251	S-199	8.0	P-42	112000
252	S-21 1	0.8	P-42	<b>245</b> 0
253	S-22 2	0.8	P-42	800
254	S-233	0.8	P-42	3250
255	<b>S-26</b> 5	0.8	P-42	43000
256	S-27 7	0.8	P-42	37000
257	<b>S-2</b> 8 3	0.8	P-42	71000
258	S-297	0.8	P-42	5800
259	S-30 :	0.8	P-42	375
260	S-31)	0.8	P-42	. 650
261	S-32 ?	0.8	P-42	2400
262	S-17 7	0.8	P-46	68000
263	<b>S-18</b> 3	0.8	P-46	13000
264	S-19 )	0.8	P-46 P-46	88000 2900
265	S-21 1	0.8 0.8	P-46	2900 1400
266	S-22 2		P-46	
<b>267</b>	S-23 3	0.8	<b>1</b> -40	2400

Table U
(Continued)

		Surfactar		
		Concentr		Brookfield Viscosity
<b>Example</b>	<u>Surfaactant</u>	(wt.%)	<u>Thickener</u>	(cps) @ pH = 9.0
<b>26</b> 8	S-266	0.8	P-46	25000
269	S-227	0.8	P-46	38500
270	S-288	0.8	P-46	77000
271	S-229	0.8	P-46	7200
272	S-300	0.8	P-46	<b>55</b> 0
273	S-3B1	0.8	P-46	<b>69</b> 0
274	S-322	0.8	P-46	1775
275	Aerrosol C	O.0 TC	P-4	50500
	Aerrosol C	OT 0.1	P-4	93500
	Aerrosol C	OT 0.2	P-4	<b>42000</b>
	Aerrosol C	T 0.4	P-4	11200
	Aerrosol C	0.8 TC	P-4	3700
	Aerrosol C	OT 1.0	P-4	<b>720</b> 0
	Aerrosol C	OT 2.0	P-4	10600

#### Examples 276-288

## Co-TThickening with Solvents and Non-Solvents

Solvents and non-solvents added to an associative polymer solution influences the co-thickening effect. The results in Table W below show the co-thickening effect produced by the addition with thorough mixing of certain solvents and non-solvents identified in Table V below in the amounts identified in Table W to a 0.75% alkaline solution of an alkali soluble thickener identified in Table W as measured with a Brookfield Viscometer at 6 rpm at pH = 9.0.

- 95 -

## Table V

Solvwent Desisignation	Solvent
O-1 . O-2 ! O-3 ! O-4 : O-5 : O-6 : O-7 ' O-8 : O-9 !	mineral spirits butanol Isobutanol Isopropanol 2-Ethylhexanol Butyl Carbitol Butyl DiPropasol Butyl Propasol Propyl DiPropasol Propyl Propasol
O-111 O-122	Methyl DiPropasol Methyl Propasol
V	1/2001/J1 2 10/paio01

## Table W

Example	Thickenerer	Solvent	Solvent Concentration(wt.%)	Solvent O-1 Concentration (wt.%)	Brookfield Viscosity (cps) @ pH = 9,0
276	P-3	O-6	5	0	29200
	P-3	O-6	10	0	865
	P-3	0-6	20	0	625
	P-3	0-6	40	0	720
	P-3	O-6	5	5	15400
	P-3	O-6	10	5	1125
	P-3	O-6	20	5	<b>73</b> 5
	P-3	O-6	40	5	780
	P-3	O-6	5	10	56500
	P-3	O-6	10	10	1050
	P-3	O-6	20	10	835
	P-3	O-6	40	10	832
	P-3	O-6	5	20	41500
	P-3	O-6	10	· <b>20</b>	1625

Table W (Continued)

Example	Thickenmer	Solvent	Solvent Concentration (wt.%)	Solvent O-1 Concentration (wt.%)	Brookfield Viscosity (cps) @ pH = 9.0
277	P-3	0-7	. 0	0	76000
	P-3	0-7	5	0	2150
	P-3	0-7	10	0	3700
	P-3	0-7	20	0	2000
	P-3	0-7	0	5	89000
	P-3	0-7	5	5	88000
	P-3	0-7	.10	5	50000
	P-3	0-7	20	5	46500
	P-3	0-7	0	10	102400
	P-3	0-7	5	10	122000
	P-3	0-7	10	10	72000
	P-3	0-7	0	20	113000
	P-3	0-7	5	20	158000
	P-3	O-7	10	20	138000
278	P-3	O-8	5	0	1925
	P-3	O-8	10	0	1150
	P-3	O-8	20	. 0	2000
	P-3	O-8	40	0	6200
279	P-3	<b>O</b> -9	5	0	36000
	P-3	O-9	10	0	1200
	P-3	O-9	20	0	<b>44</b> 0
	P-3	O-9	40	0	<b>137</b> 5
280	P-3	0-10	5	0	1375
	P-3	0-10	10	0	45500
	P-3	0-10	20	0	<b>62</b> 5
	P-3	0-10	40	0	510
281	P-3	0-11	5	0	36000
•	P-3	0-11	10	0	20500
	P-3	0-11	20	0	4200
	P-3	0-11	40	0 ·	550

. . J. YAY YM. KULL

<u>Table W</u> (Continued)

Example	Thickener r	Solvent	Solvent Concentration (wt.%)	Solvent O-1 Concentration (wt%)	Brookfield Viscosity (cps) @ pH = 9.0
282	P-3	0-12	0	0	76000
	P-3	0-12	5	0	45000
	P-3	0-12	10	0	24500
	P-3	0-12	20	0	5800
	P-3	0-12	40	0	<b>67</b> 5
	P-3	0-12	5	5	51500
	P-3	0-12	10	5	28500
	P-3	0-12	20	5	7100
	P-3	0-12	40	5	810
	P-3	0-12	5	10	61200
	P-3	0-12	10	10	33500
	P-3	0-12	20	10	6400
	P-3	0-12	40	10	950
	P-3	0-12	5	20	86800
	P-3	0-12	10	20	40500
	P-3	0-12	20	20	7100
	P-3	O-12	40	20	1350
283	P-14	0-7	0	0	150000
	P-14	O-7	5	0	1350
	P-14	0-7	10	0	4500
	P-14	0-7	20	0	7000
	P-14	0-7	0	5	140000
	P-14	0-7	5	5	120000
	P-14	0-7	10	5	78000
	P-14	0-7	0	5	<b>1400</b> 00
	P-14	0-7	5	10	158000
	P-14	O-7	10	10	<b>124000</b>
	P-14	O-7	0	20	136000
	P-14	O-7	5	20	15 <b>200</b> 0
	P-14	0-7	10	20	142000

<u>Table W</u> (Continued)

Example	Thickeneter	Solvent	Solvent Concentration (wt.%)	Solvent O-1 Concentration (wt%)	Brookfield Viscosity (cps) @ pH = 9.0
284	P-3a	0-2	. 0	0	132600
	P-3a	0-2	5	0	17300
	P-3a	0-2	10	0	850
	P-3a	0-2	20	0	1425
	P-3a	0-2	40	0	4750
	P-3a	0-2	0	5	140000
	P-3a	0-2	5	5	<b>6700</b> 0
	P-3a	0-2	10	5	<b>250</b> 0
	P-3a	0-2	20	5	3000
	P-3a	0-2	0	10	134000
	P-3a	0-2	5	10	33000
	P-3a	0-2	10	10	4000
	P-3a	0-2	20	10	4900
	P-3a	0-2	0	20	144000
	P-3a	0-2	5	20	49000
	P-3a	O-2	10	20	8000
285	P-3a	O-3	5	0	28500
	P-3a	O-3	10	0	880
	P-3a	O-3	20	0	1425
	P-3a	O-3	40	0	4600
	P-3a	O-3	5	5	80000
	P-3a	O-3	10	5	<b>295</b> 0
	P-3a	O-3	20	5	3200
	P-3a	O-3	40	5	6200
	P-3a	0-3	5	10	78000
	P-3a	O-3	10	10	5200
	P-3a	O-3	20	10	6400
	P-3a	O-3	5	<b>2</b> 0	136000
	P-3a	O-3	10	20	20500

WQ 53/44544

Table W (Continued)

	•		Solvent	Solvent O-1	Brookfield
			Concentration	Concentration	Viscosity
Example	Thickenerer	Solvent	(wt.%)	(wt.%)	(cps) @ pH = 9.0
286	P-3a	0-4	5	0	94000
	P-3a	0-4	10	0	29000
	P-3a	0-4	20	0	1050
	P-3a	0-4	40	0	850
	P-3a	0-4	5	5	107400
	P-3a	0-4	10	5	39000
	P-3a	0-4	20	5	1225
	P-3a	0-4	40	5	900
	P-3a	0-4	5	10	134000
	P-3a	0-4	10	10	41000
	P-3a	0-4	20	10	1350
	P-3a	0-4	40	10	1050
	P-3a	0-4	5	20	164000
	P-3a	0-4	10	20	33000
	P-3a	0-4	20	20	1825
	P-3a	0-4	40	20	<b>135</b> 0
287	P-3a	O-5	5	0	93500
	P-3a	O-5	10	0	136000
	P-3a	0.5	20	0	178000
288	P-3a	0-7	5	0	2700
	P-3a	0-7	10	0	6100
	P-3a	0-7	20	0	11900

Although the invention has been illustrated by certain of the preceding examples, it is not to be construed as being limited thereby; but rathoer, the invention encompasses the generic area as hereinbefore discklosed. Various modifications and embodiments can be made without a departing from the spirit and scope thereof.

- 100 -

#### Claims

- 1. A polymer comprising the reaction product of:
- (A)) about 0-99.9 weight percent of one or more nonionic, cationnic, anionic or amphoteric monomers;
- (B)) about 0-99.9 weight percent of one or more monoethylenically unsaturated monomers different from (A);
- (C)) about 0.1-100 weight percent of one or more monoethylenically unsaturated macromonomers different from (A) and (B); and
- (D)) about 0-20 weight percent or greater of one or more polyethylenicallyly unsaturated monomers different from (A), (B) and (C).
- 2.2. The polymer of claim 1 comprising the reaction product of:
- (AA) about 1-99.9 weight percent of one or more alpha, beta-monoethyldenically unsaturated carboxylic acids;
- (EB) about 0-98.9 weight percent of one or more monoethylenically unsaturated monomers different from (A);
- (CC) about 0.1-99 weight percent of one or more monoethylenically unsaturated macromonomers different from (A) and (B); and
- (ED) about 0-20 weight percent or greater of one or more polyethyleenically unsaturated monomers different from (A), (B) and (C).
- 3.3. The polymer of claim 1 comprising the reaction product of:
- (AA) from 1 to about 99.9 weight percent of one or more nonionic nmonoethylenically unsaturated monomers; and

- 101 -

- (B);) from about 0.1 to 99 weight percent of one or more monoethyleenically unsaturated macromonomers different from (A).
  - 4. A compound represented by the formula:

$$\begin{array}{c} RR^4\\ \text{II}\\ R^1\text{-}(OR^2)_z\text{-}R^3\text{-}C\text{-}\text{=}CR^5R^6 \end{array}$$

wherein:

R<sup>11</sup> is a monovalent residue of a substituted or unsubstituted complex hydrophobe compound;

each  $R^2$  is the same or different and is a substituted or unsubstituted divvalent hydrocarbon residue;

 $\mathbb{R}^{3\,3}$  is a substituted or unsubstituted divalent hydrocarbon residdue;

 ${\bf R^4, i}$ ,  ${\bf R^5}$  and  ${\bf R^6}$  are the same or different and are hydrogen or a substituted or unsubstituted monovalent hydrocarbon residue; and

z is s a value of 0 or greater.

5. A compound represented by the formula selected from:

wherein  $R_1$  and  $l R_2$  are the same or different and are hydrogen or a substituted or unnsubstituted monovalent hydrocarbon residue,  $R_3$  is a substituted or unnsubstituted divalent or trivalent hydrocarbon residue, each  $R_{44}$  is the same or different and is a substituted or unsubstituted difivalent hydrocarbon residue, each  $R_5$  is the same or different and is an substituted or unsubstituted divalent hydrocarbon residue, R<sub>6</sub> is hyydrogen, a substituted or unsubstituted monovalent hydrocarbon resisidue or an ionic substituent, a and b are the same or different and aree a value of 0 or 1, and x and y are the same or different and are e a value of 0 or greater; provided (i) at least two of R<sub>1</sub>,  $R_2$ ,  $R_3$ ,  $R_4$ ,  $R_5$  aand  $R_6$  are a hydrocarbon residue having greater than 2 carbon atoms in the case of  $R_1$ ,  $R_2$  and  $R_6$  or having greater than 2 pendant carbon s atoms in the case of  $R_3$ ,  $R_4$  and  $R_5$ , (ii) at least one of  $R_4$  and  $R_5$  is other than -CH<sub>2</sub>CH<sub>2</sub>- when  $R_1$  and  $R_2$  are nonylphenyl, R<sub>3</sub> is CH-, R<sub>6</sub> is s hydrogen or SO<sub>3</sub>, a is a value of 1, b is a value of 1, and at least one  $\epsilon$  of x and y is a value of 1 or greater, and (iii)  $R_A$  is -CH<sub>2</sub>CH<sub>2</sub>- and xx is a value of at least 2 when  $R_1$  is hexadecyl,  $R_2$  is tetradecyl, R<sub>3</sub> is CH-, at least one R<sub>5</sub> is -CH<sub>2</sub>CH(tetradecyl)-, R<sub>6</sub> is hydrogen, y is a value of 1 or greater, a is a value of 1, and b is a value of 0; and

wherein  $R_7$  and  $1\,R_8$  are the same or different and are hydrogen or a substituted or unnsubstituted monovalent hydrocarbon residue,  $R_9$  and  $R_{12}$  are the : same or different and are a substituted or

PCT/US93/04872

VO 93/24544

unsubstituted divialent or trivalent hydrocarbon residue, each  $R_{10}$  is the same or different and is a substituted or unsubstituted divalent hydrocarbon residue, each  $R_{13}$  is the same or different and is a substituted or unnsubstituted divalent hydrocarbon residue,  $R_{11}$  and  $R_{14}$  are the samee or different and are hydrogen, a substituted or unsubstituted monovalent hydrocarbon residue or an ionic substituted,  $R_{15}$  i is a substituted or unsubstituted divalent hydrocarbon residue, d and e are the same or different and are a value of 0 or 1, annd f and g are the same or different and are a value of 0 or greater; provivided at least two of  $R_7$ ,  $R_8$ ,  $R_9$ ,  $R_{10}$ ,  $R_{11}$ ,  $R_{12}$ ,  $R_{13}$ ,  $R_{14}$  and  $R_{15}$  are a hydrocarbon residue having greater than 2 carbon atoms in the case of  $R_7$ ,  $R_8$ ,  $R_{11}$  an  $R_{14}$  or having greater than 2 pendant carbon automs in the case of  $R_9$ ,  $R_{10}$ ,  $R_{12}$ ,  $R_{13}$  and  $R_{15}$ .

- 6. The polymers of claims 1, 2 and 3 wherein said monoethylenically unsaturated macromonomer is represented by the formula depicted 1 in claim 4.
- 7. The polymers of claim 6 wherein the substituted or unsubstituted complex hydrophobe compound is represented by the formula selected 1 from:

wherein  $R_1$  and  $IR_2$  are the same or different and are hydrogen or a substituted or unnsubstituted monovalent hydrocarbon residue,  $R_3$  is a substituted or unnsubstituted divalent or trivalent hydrocarbon residue,

- 104 -

each  $R_4$  is the ssame or different and is a substituted or unsubstituted divalent hydroccarbon residue, each  $R_5$  is the same or different and is a substituted or unsubstituted divalent hydrocarbon residue,  $R_6$  is hydrogen, a substituted or unsubstituted monovalent hydrocarbon residue or an ionic substitutent, a and b are the same or different and are a value of 0 or 1, and x s and y are the same or different and are a value of 0 or greater; provideed at least two of  $R_1$ ,  $R_2$ ,  $R_3$ ,  $R_4$ ,  $R_5$  and  $R_6$  are a hydrocarbon ressidue having greater than 2 carbon atoms in the case of  $R_1$ ,  $R_2$  and  $R_6$  oor having greater than 2 pendant carbon atoms in the case of  $R_3$ ,  $R_4$  and  $R_6$ ; and

 $R_{7}$ -(OCH<sub>2</sub>)<sub>d</sub>- $R_{9}$ -)-(OR<sub>10</sub>)<sub>f</sub>-OR<sub>11</sub> |  $R_{1515}$ |  $R_{8}$ -(OCH<sub>2</sub>)<sub>e</sub>- $R_{122}$ -(OR<sub>13</sub>)<sub>g</sub>-OR<sub>14</sub>

wherein  $R_7$  and  $d_8$  are the same or different and are hydrogen or a substituted or unsubstituted monovalent hydrocarbon residue,  $R_9$  and  $R_{12}$  are the same or  $d_8$  different and are a substituted or unsubstituted divalent or trivalent hydrocarbon residue, each  $R_{10}$  is the same or different and is a substituted or unsubstituted divalent hydrocarbon residue, each  $R_{13}$  is the same or different and is a substituted or unsubstituted divalent hydrocarbon residue,  $R_{11}$  and  $R_{14}$  are the same or different and are hydrogen, a substituted or unsubstituted monovalent hydrocarbon residue or an ionic substituted,  $R_{15}$  is a substituted or unsubstituted divalent hydrocarbon residue, d and e are the same or different and are a value of 0 or 1, and f and gg are the same or different and are a value of 0 or greater; provided at least two of  $R_7$ ,  $R_8$ ,  $R_9$ ,  $R_{10}$ ,  $R_{11}$ ,  $R_{12}$ ,  $R_{13}$ ,  $R_{14}$  and  $R_{15}$  are a hydrocarbon residue having greater than 2 carbon atoms in the case of

- 105 -

 $\rm R_7, R_8, R_{11}$  an  $\rm RR_{14}$  or having greater than 2 pendant carbon atoms in the case of  $\rm R_9, R_{10}, RR_{12}, R_{13}$  and  $\rm R_{15}.$ 

8. The compound of claim 4 wherein the substituted or unsubstituted complex hydrophobe compound is represented by the formula selected 1 from:

wherein  $R_1$  and  $FR_2$  are the same or different and are hydrogen or a substituted or unasubstituted monovalent hydrocarbon residue,  $R_3$  is a substituted or unnsubstituted divalent or trivalent hydrocarbon residue, each  $R_A$  is the samme or different and is a substituted or unsubstituted divalent hydrocarrbon residue, each R5 is the same or different and is a substituted or unnsubstituted divalent hydrocarbon residue,  $R_{c}$  is hydrogen, a substitiuted or unsubstituted monovalent hydrocarbon residue or an ionic substitituent, a and b are the same or different and are a value of 0 or 1, and x annd y are the same or different and are a value of 0 or greater; provided ( (i) at least two of  $R_1$ ,  $R_2$ ,  $R_3$ ,  $R_4$ ,  $R_5$  and  $R_6$  are a hydrocarbon residdue having greater than 2 carbon atoms in the case of R<sub>1</sub>, R<sub>2</sub> and R<sub>6</sub> or r having greater than 2 pendant carbon atoms in the case of  $R_3$ ,  $R_4$  and  $R_5$ , , (ii) when  $R_1$  is hexadecyl, a is a value of 1,  $R_2$  is tetradecyl, b is a vvalue of 0, R<sub>3</sub> is CH-, R<sub>4</sub> is -CH<sub>2</sub>CH(tetradecyl)-, x is a value of 1, R<sub>5</sub> is -CCH<sub>2</sub>CH<sub>2</sub>-, y is a value of 34, R<sub>6</sub> is hydrogen and z is a value of 0, then thhe  $-R^3$ - $(R^4)$ C= $CR^5R^6$  portion of the compound is other than the residue ( of maleic anhydride, and (iii) when  $R_1$  is hexadecyl, a is a value of 1,  $R_2$  is 3 tetradecyl, b is a value of 0,  $R_3$  is CH-,  $R_4$  is

- 106 -

-CH<sub>2</sub>CH(tetradelecyl)-, x is a value of 1, y is a value of 0,  $R_6$  is hydrogen and z is a value of 344, then the -R<sup>3</sup>-(R<sup>4</sup>)C=CR<sup>5</sup>R<sup>6</sup> portion of the compound is other than the residue of maleic anhydride; and

$$R_{7}$$
-(OCH<sub>2</sub>)<sub>d</sub>- $R_{9}$ -(-(OR<sub>10</sub>)<sub>f</sub>-OR<sub>11</sub> | R<sub>155</sub> | R<sub>8</sub>-(OCH<sub>2</sub>)<sub>e</sub>- $R_{122}$ -(OR<sub>13</sub>)<sub>e</sub>-OR<sub>14</sub>

wherein  $R_7$  and  $1R_8$  are the same or different and are hydrogen or a substituted or unsubstituted monovalent hydrocarbon residue,  $R_9$  and  $R_{12}$  are the same or  $\epsilon$  different and are a substituted or unsubstituted divalent or trivalent hydrocarbon residue, each  $R_{10}$  is the same or different and is a substituted or r unsubstituted divalent hydrocarbon residue, each  $R_{13}$  is the same or different and is a substituted or unsubstituted divalent hydrocarbon residue,  $R_{11}$  and  $R_{14}$  are the same or different and are hydrogen, a substituted or unsubstituted monovalent hydrocarbon residue or an ionic substituted or unsubstituted or unsubstituted divalent hydrocarbon residue, d and e are the same or different and are a value of 0 or 1, and f and gg are the same or different and are a value of 0 or greater; provided at leastst two of  $R_7$ ,  $R_8$ ,  $R_9$ ,  $R_{10}$ ,  $R_{11}$ ,  $R_{12}$ ,  $R_{13}$ ,  $R_{14}$  and  $R_{15}$  are a hydrocarbon ressidue having greater than 2 carbon atoms in the case of  $R_7$ ,  $R_8$ ,  $R_{11}$  an I  $R_{14}$  or having greater than 2 pendant carbon atoms in the case of  $R_9$ ,  $R_{10}$ ,  $R_{10}$ ,  $R_{12}$ ,  $R_{13}$  and  $R_{15}$ .

9. The polymers of claim 7 and the compound of claim 8 wherein a (i) at least one of R<sub>1</sub>, R<sub>2</sub>, R<sub>7</sub> and R<sub>8</sub> is a hydrocarbon radical represented by the formula:

4,U 3312-13-14

- 107 -

wherein  $R_{16}$  and  $d_{17}$  are the same or different and are hydrogen or a substituted or unnsubstituted monovalent hydrocarbon residue,  $R_{18}$  is a substituted or unnsubstituted divalent or trivalent hydrocarbon residue, and h annd i are the same or different and are a value of 0 or 1; and/or (ii) at lexast one of  $R_4$ ,  $R_5$ ,  $R_{10}$  and  $R_{13}$  is a hydrocarbon radical representated by the formula:

# $-CHH[(OR_{19})_jOR_{20}]-$

wherein each R<sub>199</sub> is the same or different and is a substituted or unsubstituted divivalent hydrocarbon residue, R<sub>20</sub> is hydrogen; a substituted or unnsubstituted monovalent hydrocarbon residue or an ionic substituent,; and j is a value of 0 or greater.

10. The polymers of claim 7 and the compound of claim 8 in which 1 said monoethylenically unsaturated macromonomer and compound are represented by the formula selected from:

ママン せいにんりゅん

- 108 -

$$\begin{array}{c} R^4\\ \mid\\ R^1\text{-}(OR^2)_z\text{-}CH_2CCHCH_2OCH_2C=CH_2\\ \mid\mid\\ (OOR_{19})_jOH \end{array}$$

$$R^4$$

$$R^1 - (OR^2)_z - OCC(O) - C = CH_2$$

wherein  $R^1$ ,  $R^2$ ,  $R^4$  and  $R^4$  and  $R^4$  are as defined in claim 4, and  $R^4$  and  $R^4$  are as defined in claim 9.

- 11. An aqueous latex containing the solubilized or neutralized aqueeous polymer of claim 1.
- 12. A process of thickening an aqueous system comprising addining to the system a polymer of claim 1 and at least partially solubilizing or neutralizing said polymer.
- 13. The process of claim 12 wherein the system is thickened furtheer by the addition of an effective amount of surfactant, solvent or non-scolvent.
- 14. An aqueous polymer emulsion useful as a thickening agentt in aqueous compositions when the polymer is

- 109 -

dissolved thereing, which comprises water and emulsified polymer particles comprisising the polymer of claims 1, 2 or 3.

- polymer, water, ppigment, and thickener, the improvement which comprises having g as at least a portion of the thickener dissolved therein a polymer comprising the polymer of claims 1, 2 or 3.
- 16. In an aqueous composition comprising water and thickener, the improvement which comprises having as at least a portion of the thickener dissolved therein a polymer comprising the polymer of claims s 1, 2 or 3.
  - 17. A derivative of the compound of claims 4 or 5.
- 18. An emulsifier, dispersant, stabilizer, wetting agent, a copolymeerizable monomer in polymerization reaction or a latex binder compprising the compound of claims 4 or 5.
- 19. A latex composition containing water, latex polymer and the c compound of claims 4 or 5.
- 20. A method for the alkoxylation of a complex hydrophobe compound having at least one active hydrogen comprising alkoxyylating the complex hydrophobe compound of claim 5 with an alkylence oxide in the presence of a catalyst to produce alkoxylates of thee complex hydrophobe compound under alkoxylation conditions.
- 21. A method for preparing alkoxylation product mixtures having a a narrow distribution of species with at least one specie comprising at least about 10 weight percent of the product

WO 93/24544

PCT/US93/04872

-110 -

mixture compprising contacting a complex hydrophobe compound of claim 5 havings at least one active hydrogen with an alkylene oxide in the presence oxf a catalytically effective amount of catalyst under alkoxylation cconditions sufficient to provide a product mixture having at least one alkoxylation specie comprising at least about 10 weight percent of the mixture.

222. The method of claim 20 further comprising derivatizing the product in which the derivatizing reaction comprises an oxidation, c carboxylation, reduction, hydrogenation, dehydrogenation, condensation, amination, esterification, etherification, alkylation or acylation reaction.

3 1

A. CLASSIFICATION OF SUBJECT MATTITER

IPC 5 CO8F220/28 C08BF220/04 C08F246/00 C09D4/00 According to International Patent Classificationon (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classificatiation system followed by classification symbols) IPC 5 COSF CO9D CO7C: Documentation searched other than minimum to documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the interprnational search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE REFLEVANT Category \* Citation of document, with indicatation, where appropriate, of the relevant passages Relevant to claim No. X US,A,4 029 872 ((EVANI ET AL.) 14 June 1977 1,3,4,6, 16-18 see the whole dalocument X EP, A, 0 173 109 ((DESOTO) 5 March 1986 1-8. 10-19 cited in the appplication see claims 1-10) & US, A, 4 514 5522 X EP,A,O 222 059 ((DIAMOND SHAMROCK 1,4,5 CHEMICALS) 20 Malay 1987 see page 5, linee 1 - page 6, line 40 X EP,A,O 216 479 ((ALLIED COLLOIDS) 1 April 1-4,6, 1987 11-13, 16-18 see claim 1 -/--X Further documents are listed in the continuation of box C. Patent family members are listed in sonex. Special categories of cited documents: T later document published after the international filing date or priority date and not in conflict with the application bu-cited to understand the principle or theory underlying the "A" document defining the general state of the se art which is not considered to be of particular relevance "E" earlier document but published on or after to the international "X" document of particular relevance; the elained invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date "L" document which may throw doubts on priorierly claim(s) or which is cited to establish the publication or date of another citation or other special reason (as specificfied) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, us use, exhibition or ments, such combination being obvious to a person skilled other means "P" document published prior to the internationonal filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the internationanal search Date of mailing of the international search report 1 1. 11. 93 4 November 1993 Name and mailing address of the ISA **Authorized officer** European Patent Office, P.B. 5( 5818 Patendaan 2 NL - 2220 HV Rijewijk Tel. (+31-70) 340-2040, Tz. 3131 651 epo nl, Fax (+31-70) 340-3016

Form PCT/ISA/210 (second sheet) (July 1992)

2

ANDRIOLLO, G

		PCT/US 93/04872
C.(Continue	DOCUMENTS CONSIDERRED TO BE RELEVANT	
Category *	Citation of document, with indicalestion, where appropriate, of the relevant passages	Relevant to claim No.
X	EP,A,O 011 806 ((DDW CHEMICAL) 11 June 1980 cited in the appplication see claims 1-111 & US,A,4 384 0996	1-4,6, 11-13, 16-18
- - - - - - - - - - - - - - - - - - -		
	·	
		×
		·

# IN NATIONAL SEARCH REPORT

In the least Application No PC-1/US 93/04872

		PUTY	US 93/U48/Z
Patent document cited in search report	Publication date	Patent family member(s)	Publication date
US-A-4029872	14-06-77	US-A- 3963684 US-A- 4038265 US-A- 4029874 US-A- 4029873	26-07-77
EP-A-0173109	05-03-86	US-A- 4514552 AU-B- 573021 AU-A- 4613685 CA-A- 1278893 JP-A- 61060719 US-E- RE33156	30-04-85 26-05-88 27-02-86 08-01-91 28-03-86 30-01-90
US-A-4514552	30-04-85	AU-B- 573021 AU-A- 4613685 CA-A- 1278893 EP-A,B 0173109 JP-A- 61060719 US-E- RE33156	26-05-88 27-02-86 08-01-91 05-03-86 28-03-86 30-01-90
EP-A-0222059	20-05-87	US-A- 4876384 AU-B- 595219 AU-A- 6041986 CA-A- 1292476 DE-A- 3685938 JP-A- 62114932 US-A- 5053554 US-A- 5110889 US-A- 5159119 US-A- 5243085	24-10-89 29-03-90 21-05-87 26-11-91 13-08-92 26-05-87 01-10-91 05-05-92 27-10-92 07-09-93
EP-A-0216479	01-04-87	AU-A- 4621785 AU-B- 612965 AU-A- 6089786 CA-A- 1241492 CA-A- 1318689 CA-A- 1264280 CA-A- 1295778 DE-A- 3584551 EP-A,B 0172723 EP-A,B 0172724	20-02-86 25-07-91 19-02-87 30-08-88 01-06-93 09-01-90 11-02-92 05-12-91 26-02-86 26-02-86

ional Application No PCT/US 93/04872

			F01703	33/0 <del>10</del> /2
Patent document cited in search report	Publication date	Patent fa member	mily r(s)	Publication date
EP-A-0216479		EP-A,B	0172025	19-02-86
		EP-A,B	0213799	11-03-87
		EP-A,B	0213800	11-03-87
		EP-A,B	0215565	25-03-87
		EP-A,B	0214760	18-03-87
		JP-A-	62042731	<b>24-02-87</b> .
			62039537	20-02-87
			62057408	13-03-87
		JP-A- (	62039608	20-02-87
		US-A-	4892916	09-01-90
		US-A-	4940763	10-07-90
		US-A-	4980434	25-12-90
		US-A-	5210324	11-05-93
	•	US-A-	4702844	27-10-87
		us-a-	4792343	20-12-88
		US-A-	4741790	03-05-88
EP-A-0011806	11-06-80	AT-T-	5415	15-12-83
	-	AU-B-	527072	17-02-83
		AU-A-	5295379	29-05-80
	•	CA-A-	1152668	23-08-83
	•	US-A-	4384096	17-05-83
	• •	JP-C-	1486672	14-03-89
			55073747	03-06-80
		JP-B- 6	3035672	15-07-88
US-A-4384096	17-05-83	AT-T-	5415	15-12-83
		AU-B-	527072	17-02-83
		AU-A-	5295379	29-05-80
		CA-A-	1152668	23-08-83
		EP-A,B	0011806	11-06-80